



App No.: 10/764,390 Docket No.: 511582008100  
Inventor: Steven B. KANNER et al.  
Title: NUCLEIC ACIDS AND CORRESPONDING PROTEINS  
ENTITLED 254P1D6B USEFUL IN TREATMENT AND  
DETECTION OF CANCER

Sheet 1 of 38

Figure 1: 254P1D6B SSH sequence of 186 nucleotides (SEQ ID NO: 1).

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1 GATCCACAGA TAGGACACAA TTCTTGGTC ATCAGTAGAC CTTGAACCAT CCAAAGTAAT
61 GGAATTATTG GGAAGCACAA GAACATGTCT GCCACCAGCC CGGGCTCTGG GAGGACTATT
121 ATTTCCCTTC TTCACAGCCA CAGTGAGGGT GGACGTGCTG CTCAGTCCCT GCTGGTCTTT
181 TACTGTAAA CGGAAGTGGT AGGTCCCCAC CTGGAGACCA GTCACAGTGG CTATTGCTTT
241 GTCAATATTT TCCATCTCCA CTGCACTGGG GCCTCTGACG TGCT
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**Figure 2:**

**Figure 2A. The cDNA (SEQ ID NO. : 2) and amino acid sequence (SEQ ID NO. : 3) of 254P1D6B v.1 clone LCP-3.**  
The start methionine is underlined. The open reading frame extends from nucleic acid 512-3730 including the stop codon.

1 gctgccgcggcggtggcgggatccccggggatggcaacccgtccacccgtgc  
61 cctcggcgccctggctggcccgccagagcggcgccgctcgactgcccga  
121 ggtgagagcgcagcagtagttcagcctgtcttggcttccagattcgctcccttgg  
181 ggctacgtcccgaaaaagaggaagcggaggatttgcgtgggtgggggttacctttaac  
241 agcagggtgcgcgcgcgagggtgttacgtgtgtgtgtgtgtgtgtgtgt  
301 taagacctgcgtatgcgtacgacgaggagaacaagtggacggcgtatgtcaggccag  
361 cagcaacgcgtggcgagcttcagtgccacgcgtgaccacagtttgcaggccaaa  
421 tctggctctaaaaacatcaaaggaaagcttgcaccaaactcttcaggccgtcag  
1 M A P P T G V L S S  
481 aagcctgcgtatgcgttgcacaATGGCGCCCCCACAGGTGTGCTCTCTTC  
11 L L L L V T I A G C A R K Q C S E G R T  
541 ATTGCTGCTGCTGGTACAATTGCAGGTTGTGCCGTAAGCAGTGCAGCGAGGGAGGAC  
31 Y S N A V I S P N L E T T R I M R V S H  
601 ATATTCCAATGCAGTCATTCACCTAACCTGGAAACCACAGAATCATGCGGGTGTCTCA  
51 T F P V V D C T A A C C D L S S C D L A  
661 CACCTTCCCTGCGTAGACTGCACGGCCGCTTGCTGTGACCTGTCCAGCTGTGACCTGGC  
71 W W F E G R C Y L V S C P H K E N C E P  
721 CTGGTGGTTCGAGGGCCGCTGCTACCTGGTGAAGCTGCCAACAAAGAGAACTGTGAGCC  
91 K K M G P I R S Y L T F V L R P V Q R P  
781 CAAGAAGATGGGCCCATCAGGTCTTATCTCACTTTGTGCTCCGGCTGTTCAAGAGGC  
111 A Q L L D Y G D M M L N R G S P S G I W  
841 TGCACAGCTGGACTATGGGACATGATGCTGAACAGGGCTCCCCCTGGGATCTG  
131 G D S P E D I R K D L P F L G K D W G L  
901 GGGGACTCACCTGAGGATATCAGAAAGGACTTGMCCCTTCTAGGCAAAGATTGGGCCT  
151 E E M S E Y S D D Y R E L E K D L L Q P  
961 AGAGGAGATGTCTGAGTACKCAGATGACTACCGGGAGCTGGAGAAGGACCTTTGCAACC  
171 S G K Q E P R G S A E Y T D W G L L P G  
1021 CAGTGGCAAGCAGGAGCCAGAGGGAGTGCCTGAGTACACGGACTGGGCCTACTGCCGGG  
191 S E G A F N S S V G D S P A V P A E T Q  
1081 CAGCGAGGGGGCCTCAACTCCTGTGGAGACAGTCCTGCGGTGCCAGCGGAGACGCA  
211 Q D P E L H Y L N E S A S T P A P K L P  
1141 GCAGGACCCCTGAGCTCCATTACCTGAATGAGTCGGCTCAACCCCTGCCAAAACCTCCC  
231 E R S V L L P L P T T P S S G E V L E K  
1201 TGAGAGAAGTGTGTTGCTCCCTGCCGACTACTCCATCTCAGGAGAGGTGGAGAA  
251 E K A S Q L Q E Q S S N S S G K E V L M  
1261 AGAAAAGGCTTCTCAGCTCCAGGAACAATCCAGCAACAGCTCTGGAAAAGAGGTTCTAAT  
271 P S H S L P P A S L E L S S V T V E K S  
1321 GCCTTCCCATAGTCTTCCCTCCGGCAAGCCTGGAGCTCAGTCAGTCACCGTGGAGAAAAG

291 P V L T V T P G S T E H S I P T P P T S  
 1381 CCCAGTGCTCACAGTCACCCGGGAGTACAGAGCACAGCATCCAAACACCTCCACTAG  
 311 A A P S E S T P S E L P I S P T T A P R  
 1441 CGCAGCCCCCTCTGAGTCCACCCCATCTGAGCTACCCATATCTCCTACCACTGCTCCAG  
 331 T V K E L T V S A G D N L I I T L P D N  
 1501 GACAGTGAAAGAACTTACGGTATCGGCTGGAGATAACCTAATTATAACTTACCCGACAA  
 351 E V E L K A F V A P A P P V E T T Y N Y  
 1561 TGAAGTTGAAGTGAAGGCCCTTGTTGCGCCAGGCCACCTGTAGAAACAAACCTACAACTA  
 371 E W N L I S H P T D Y Q G E I K Q G H K  
 1621 TGAATGGAATTAAAGCCACCCACAGACTACCAAGGTGAAATAAAACAAGGACACAA  
 391 Q T L N L S Q L S V G L Y V F K V T V S  
 1681 GCAAACCTTAACCTCTCAATTGTCGTCGGACTTTATGTCTTCAAAGTCACTGTTTC  
 411 S E N A F G E G F V N V T V K P A R R V  
 1741 TAGTGAAAACGCCCTTGAGAAGGATTGTCAATGTCAGTGTAAAGCCTGCCAGAAGAGT  
 431 N L P P V A V V S P Q L Q E L T L P L T  
 1801 CAACCTGCCACCTGTAGCAGTTCTCCCAACTGCAAGAGCTCACTTGCCCTTGAC  
 451 S A L I D G S Q S T D D T E I V S Y H W  
 1861 GTCAGCCCTATTGATGGCAGCAAAGTACAGATGATACTGAAATAGTGAGTTATCATTG  
 471 E E I N G P F I E E K T S V D S P V L R  
 1921 GGAAGAAATAAACGGGCCCTCATAGAAGAGAAGACTTCAGTTGACTCTCCGTCTTACG  
 491 L S N L D P G N Y S F R L T V T D S D G  
 1981 CTTGTCTAACCTTGATCCTGGTAACTATAGTTCAGGTTGACTGTTACAGACTCGGACGG  
 511 A T N S T T A A L I V N N A V D Y P P V  
 2041 AGCCACTAACTCTACAACTGCAGCCCTAATAGTGAACAATGCTGTGGACTACCCACCAAGT  
 531 A N A G P N H T I T L P Q N S I T L N G  
 2101 TGCTAATGCAGGGACCAAATCACACCATAACTTGCCCCAAAATCCATCACTTGAATGG  
 551 N Q S S D D H Q I V L Y E W S L G P G S  
 2161 AAACCAGAGCAGTGACGATCACCAGATTGCTCTATGAGTGGTCCCTGGGTCTGGGAG  
 571 E G K H V V M Q G V Q T P Y L H L S A M  
 2221 TGAGGGCAAACATGTGGTCATGCAGGGAGTACAGACGCCATACCTCATTATCTGCAAT  
 591 Q E G D Y T F Q L K V T D S S R Q Q S T  
 2281 GCAGGAAGGAGATTATAACATTCAGCTGAAGGTGACAGATTCTCAAGGCAACAGTCTAC  
 611 A V V T V I V Q P E N N R P P V A V A G  
 2341 TGCTGTRGTGACTGTGATTGTCAGCCTGAAAACAATAGACCTCCAGTGGCTGGCCGG  
 631 P D K E L I F P V E S A T L D G S S S S  
 2401 CCCTGATAAAAGAGCTGATCTCCCAGTGGAAAGTGCTACCCCTGGATGGAGCAGCAGCAG  
 651 D D H G I V F Y H W E H V R G P S A V E  
 2461 CGATGACCACGGCATTGTCTTCTACCACTGGGAGCACGTCAAGAGGCCAGTGCAGTGG  
 671 M E N I D K A I A T V T G L Q V G T Y H  
 2521 GATGGAAAATATTGACAAAGCAATAGCCACTGTGACTGGCTCCAGGTGGGGACCTACCA  
 691 F R L T V K D Q Q G L S S T S T L T V A  
 2581 CTTCCGTTTGACAGTGAAAGACCAGCAGGGACTGAGCAGCACGTCCACCCCTCACTGTGGC

711 V K K E N N S P P R A R A G G G R H V L V  
 2641 TGTGAAGAAGGAAAATAATAGTCCTCCCAGAGGCCGGGCTGGTGGCAGACATGTTCTTGT  
 731 L P N N S I T L D G S R S T D D Q R I V  
 2701 GCTTCCAATAATTCCATTACTTGGATGGTCAGGTCTACTGATGACCAAAGAATTGT  
 751 S Y L W I R D G Q S P A A G D V I D G S  
 2761 GTCCTATCTGTGGATCCGGATGGCCAGAGTCCAGCAGCTGGAGATGTCATCGATGGCTC  
 771 D H S V A L Q L T N L V E G V Y T F H L  
 2821 TGACCACAGTGTGGCTCTGCAGCTACGAATCTGGTGGAGGGGGTGTACACTTCCACTT  
 791 R V T D S Q G A S D T D T A T V E V Q P  
 2881 GCGAGTCACCGACAGTCAGGGGCCCTCGGACACAGACACTGCCACTGTGGAAGTGCAGCC  
 811 D P R K S G L V E L T L Q V G V G Q L T  
 2941 AGACCCTAGGAAGAGTGGCCTGGTGGAGCTGACCTGCAGGTTGGTGTGGCAGCTGAC  
 831 E Q R K D T L V R Q L A V L L N V L D S  
 3001 AGAGCAGCGGAAGGACACCCCTGTGAGGCAGCTGGCTGTGCTGCTAACGTGCTGGACTC  
 851 D I K V Q K I R A H S D L S T V I V F Y  
 3061 GGACATTAAGGTCCAGAAGATTGGGCCACTCGGATCTCAGCACCGTGATTGTGTTTA  
 871 V Q S R P P F K V L K A A E V A R N L H  
 3121 TGTACAGAGCAGGCCCTTCAAGGTTCTCAAAGCTGCTGAAGTGGCCCGAAATCTGCA  
 891 M R L S K E K A D F L L F K V L R V D T  
 3181 CATGCGGCTCTCAAAGGAGAAGGCTGACTTCTGCTTCAAGGTCTTGAGGGTTGATAC  
 911 A G C L L K C S G H G H C D P L T K R C  
 3241 AGCAGGTTGCCCTCTGAAGTGTCTGGCCATGGTCACTGCGACCCCTCACAAAGCGCTG  
 931 I C S H L W M E N L I Q R Y I W D G E S  
 3301 CATTGCTCTCACTTATGGATGGAGAACCTTATACAGCGTTATATCTGGATGGAGAGAG  
 951 N C E W S I F Y V T V L A F T L I V L T  
 3361 CAACTGTGAGTGGAGTATATTCTATGTGACAGTGTGGCTTTACTCTTATTGTGCTAAC  
 971 G G F T W L C I C C C K R Q K R T K I R  
 3421 AGGAGGTTCACTGGCTTGATCTGCTGCTGCAAAGACAAAAAGGACTAAATCAG  
 991 K K T K Y T I L D N M D E Q E R M E L R  
 3481 GAAAAAAACAAAGTACACCATCCTGGATAACATGGATGAACAGGAAAGAATGGAACGTGAG  
 1011 P K Y G I K H R S T E H N S S L M V S E  
 3541 GCCCAAATATGGTATCAAGCACCGAAGCACAGAGCACAACTCCAGCCTGATGGTATCCGA  
 1031 S E F D S D Q D T I F S R E K M E R G N  
 3601 GTCTGAGTTGACAGTGACCAGGACACAATCTCAGCCGAGAAAAGATGGAGAGAGGGAA  
 1051 P K V S M N G S I R N G A S F S Y C S K  
 3661 TCCAAAGGTTCCATGAATGGTCCATCAGAAATGGAGCTCCTTCAGTTATTGCTCAA  
 1071 D R \*  
 3721 GGACAGATAAtggcgccagttcattgtaaagtggaggaccyttgaatccargaccagtc  
 3781 agtgggagttacagcacaaaacccactcttttagaatagttcattgaccttctcccccag  
 3841 tgggttagatgttatccccacgtactaaagaccggttttgaaggcacaaaacaaaa  
 3901 ctttgcttttaactgagatgctgttaatagaataaaggctggtaaaactytaagg  
 3961 tatatacttaaaagagtttgagttttagctggcacaatctcatattaaagatgaac

4021 aacgatttcttatctgtagaaccttagagaaggtaatgaaacaaggttaaaaaggat  
4081 gatttctgtcttagcygctgtgattgcctctaaggaacacgcattctaaacacggttctc  
4141 ttgttaggacctgcagtcagatggctgttatgttaaaatagcttgcttaagaggcacggg  
4201 ccatctgtggaggtacggagtcttgcattgttagcaagcttctgtgtgcacggcaacactc  
4261 gcacagtgcacccctccctggtttaattctgtgttatgtcaatggcagtttcatct  
4321 ctctcaagaaagcagcttggccattcaagagactaaggaatacgattctaggact  
4381 gaggcaatagaaaaggggaggaggacttaatgcgcrtgcaggttgaaggtagcattgtaac  
4441 attatctttcttcttaagaaaaactacactgactcctctcggttttagcagta  
4501 tagttctctaatgtaaacrgatcccagttacattaartgcaatagaagtgattaattc  
4561 attaagcatttattatgttctgttaggctgtgcgtttggactgcctataggkataacg  
4621 actcagcaattgttatattccaaaactctgaaatacagtcagtcattactggatgg  
4681 cgtggttatgatactctggccccacaggtaattccaaaataacttgacatagatgt  
4741 ttcaattcatatgtttaaaatcatatgtttttctaccgaataatcttatttcaaa  
4801 catgaaagacaattaaacattcccccacaaagcagtactcccggcaatattactgga  
4861 gttaattgttagcctgtacgttgcggtagttcccccattccacccttggcct  
4921 gaggctggtgccctgggtgggcattttgtggaaagattagaatgagagat  
4981 agaaccagtgttggtaccaagtgtgagcacacctaaacaatattctgttgcacaatgc  
5041 ttttttaacacatggaaaacttaggaatgcattgtatgaagaagcaaggatttaaac  
5101 accagggcaggagtgcacagaaaaatgttccccatgggtcttaaaaaaaaaattcagctt  
5161 ttaggtgtttgtcatctcccgsgatttcatttcatttcattttttactt  
5221 attgtatattactggggaaaggcagaactaaaaagtgtgtcattttatatttaataat  
5281 tgcttgcttatgcctacactttctgtataactagccaaattcaataactgtctatgtt  
5341 agaaggaaaatgtgattttttttaaccagtattgagcttcataaggctagaatctg  
5401 ctttatcaggtgaccagggtatgggtttgcattgtgaaattctggcatagg  
5461 gacagcagccaaatgttaaagtcatcggcgtaatgaggaagaaggaggtaacattac  
5521 cgcttakgtacataacatgcattttacataactcatttgcatttataatcaacctt  
5581 aagaggagatactatcattttatgttgcagatagccctctgaaggcccagagaggtaa  
5641 rtaactcccgaggcatggcaagaagttagtggctccaagaactgaatgcaattttt  
5701 taaactgttagattctgtttccactaaacaaagaactcctgcctgtatggatggagg  
5761 aaattctgttggaaactttggccacctgaaagttctattccaggactaagagaattt  
5821 ctttaatggatccagagagccaaaggcagttgcagaggagatggcctgcatagtctcctgt  
5881 gatcacacccgggcccacccctcccttaggtttacagtggacttcttgccttgc  
5941 ttttctgtccttggccatctcgccttcattgcattttccatcacaaggatctt  
6001 gaatctctggaaatcaaacatcacagtagtgcattcatttttttttttttttt  
6061 ccatttctcatcagaacaaacgcacgagatggaaatgaccaaccaggcattttcatgg  
6121 ctgcatttgcatttggatcttggagataaaggcactgcataagactctggacagagaaaa  
6181 acaggccctagaatatggagttgggtttgttagggctcayargctaaacaaggacttt  
6241 ttgtctggtttacattcaatgaaggaggattcatccatggcattacaaggctaaac  
6301 tgcattactatgcacccatgcgcggggaggaggactgactaccattttttgt  
6361 aaggccatgcacccatgggtttgtgcgcggggaggaggactgactaccatttttt  
6421 gacaaagctatcatggactattttaatcttggttttattgcatttttttttttt  
6481 cctatgtgttgcacaaggatttctaatatcacactattaaatatgcactaatctaaat

6541 aaaggtgtctgtat~~ttt~~ctgtat~~ttt~~tttaggggaaattt~~ttt~~ctttatgct  
6601 tcagggttagaggattcccttgagtaggtcagcaactctggcctgcagccgtgtgt  
6661 gcacgccccatgagccgaaaagtggtcttatgtttcaa~~tgg~~ttaaaataaataaaaa  
6721 aaattt~~gaa~~acatgtgaactat~~at~~gacattcagattgtgttcataaataaagtttatt  
6781 ggaacat~~at~~cc

**Figure 2B. The cDNA (SEQ ID NO.: 4) and amino acid sequence (SEQ ID NO.: 5) of 254P1D6B v.2. The start methionine is underlined. The open reading frame extends from nucleic acid 512-3730 including the stop codon.**

1 gctccgcggcggtggcg~~gggat~~cccccg~~gggg~~tgcaac~~tt~~gctccac~~ct~~gtgc  
61 cctcggcg~~ggg~~c~~ct~~ggctggccccgc~~c~~gcagagcggcggc~~gg~~gc~~tc~~g~~ct~~gtcact~~cc~~gg  
121 ggtgagagcgc~~c~~ag~~c~~act~~tt~~cag~~c~~ctgtctggcttgc~~cc~~agattc~~g~~c~~tc~~c~~ct~~ctgg  
181 ggctacgtcccg~~ggg~~aa~~g~~ag~~g~~ga~~a~~gc~~g~~agg~~at~~tt~~g~~ctgggtggg~~ct~~gtac~~ct~~cttaac  
241 agcagg~~t~~gcgcgc~~g~~gagg~~gt~~ta~~ac~~gtgtgtgtgtgtgtgtgtgtgtgtgtgtgt  
301 taagac~~ct~~gc~~at~~gac~~g~~ac~~g~~agg~~aa~~ca~~g~~at~~gg~~ac~~gg~~gc~~ag~~tg~~at~~g~~ct~~c~~ag~~ggcc~~ag~~  
361 cagcaac~~g~~c~~at~~ggggc~~g~~ag~~ct~~c~~ag~~tg~~tc~~g~~cc~~ac~~g~~tg~~ac~~ca~~cc~~at~~ct~~ctt~~g~~agg~~cc~~aaa  
421 tctggctc~~ct~~aaaaa~~ac~~at~~ca~~aa~~g~~ga~~ag~~cttgc~~ac~~ccaa~~ac~~t~~ct~~ctt~~ca~~ggg~~cc~~c~~ct~~ca~~g~~  
1 M A P P T G V L S S  
481 aagc~~ct~~gc~~at~~ca~~cc~~act~~gt~~gt~~gg~~tc~~ac~~ATGGCGCCCCC~~AC~~AGGTGTGCTCTCTTC  
11 L L L L V T I A G C A R K Q C S E G R T  
541 ATTGCTGCTGCTGGT~~G~~ACAATTG~~C~~AGGTTGTGCC~~C~~TAAGCAGTGCAGCGAGGGAGGAC  
31 Y S N A V I S P N L E T T R I M R V S H  
601 ATATTCCAATGCAGTCATTCACCTAAC~~T~~GGAAACCACCA~~G~~AA~~T~~CATGCGGGTGTCTCA  
51 T F P V V D C T A A C C D L S S C D L A  
661 CACCTCCCTGTCG~~T~~AGACTGCACGGCCGCTTGTGACCTGTCCAGCTGTGACCTGGC  
71 W W F E G R C Y L V S C P H K E N C E P  
721 CTGGTGGTT~~C~~GAGGGCCGCTG~~T~~ACCTGGT~~G~~AGCTG~~CC~~CCCCACAAAGAGAA~~C~~TGTGAGGCC  
91 K K M G P I R S Y L T F V L R P V Q R P  
781 CAAGAAGATGGGCC~~C~~CATCAGGT~~T~~TATCTCACTTTGTGCTCCGGCTGTT~~C~~AGAGGCC  
111 A Q L L D Y G D M M L N R G S P S G I W  
841 TGCACAGCTGCTGGACTATGGGGACATGATG~~C~~TGAACAGGGCT~~CC~~CTGGGATCTG  
131 G D S P E D I R K D L P F L G K D W G L  
901 GGGGGACTCACCTGAGGATATCAGAAAGGACTTGCC~~TT~~CTAGGCAAAGATTGGGC~~CT~~  
151 E E M S E Y A D D Y R E L E K D L L Q P  
961 AGAGGAGATGTCTGAGTACGCAGATGACTACCGGGAGCTGGAGAAGGAC~~CT~~TTGCAACC  
171 S G K Q E P R G S A E Y T D W G L L P G  
1021 CAGTGGCAAGCAGGAGCC~~C~~AGAGGGAGTGCC~~G~~AGTACACGGACTGGGGCTACTGCC~~GG~~  
191 S E G A F N S S V G D S P A V P A E T Q  
1081 CAGCGAGGGGGC~~CT~~CAACTCCTGTGAGACAGTCCTGCGGTGCCAGCGGAGACGCA  
211 Q D P E L H Y L N E S A S T P A P K L P  
1141 GCAGGGACCCTGAGCTCCATTACCTGAATGAGTCGGCTTCAACCCCTGCC~~CC~~AAACTCCC  
231 E R S V L L P L P T T P S S G E V L E K

1201 TGAGAGAAAGTGTGTTGCCCTGCCGACTACTCCATCTCAGGAGAGGTGTGGAGAA  
251 E K A S Q L Q E Q S S N S S G K E V L M  
1261 AGAAAAGGCTCTCAGCTCCAGGAACAATCCAGCAACAGCTCTGGAAAAGAGGTTCTAAT  
271 P S H S L P P A S L E L S S V T V E K S  
1321 GCCTTCCCATAAGTCTTCCCTCCGGCAAGCCTGGAGCTCAGCTCAGTCACCGTGGAGAAAAG  
291 P V L T V T P G S T E H S I P T P P T S  
1381 CCCAGTGCTCACAGTCACCCCCGGGGAGTACAGAGCACAGCATCCAAACACCTCCACTAG  
311 A A P S E S T P S E L P I S P T T A P R  
1441 CGCAGCCCCCTCTGAGTCCACCCCATCTGAGCTACCCATATCTCCTACCACTGCTCCAG  
331 T V K E L T V S A G D N L I I T L P D N  
1501 GACAGTGAAGAAACTTACGGTATCGGCTGGAGATAACCTAATTATAACTTTACCCGACAA  
351 E V E L K A F V A P A P P V E T T Y N Y  
1561 TGAAGTTGAACTGAAGGCCCTTGTGCGCCAGGCCACCTGTAGAAACAAACCTACAACTA  
371 E W N L I S H P T D Y Q G E I K Q G H K  
1621 TGAATGGAATTAAATAAGCCACCCACAGACTACCAAGGTGAAATAAAACAAGGACACAA  
391 Q T L N L S Q L S V G L Y V F K V T V S  
1681 GCAAACCTTAACCTCTCAATTGTCCGTGGACTTATGTCTCAAAGTCACTGTTTC  
411 S E N A F G E G F V N V T V K P A R R V  
1741 TAGTGAACACGCCCTTGGAGAAGGATTGTCAATGTCAGTAAAGCCTGCCAGAAGAGT  
431 N L P P V A V V S P Q L Q E L T L P L T  
1801 CAACCTGCCACCTGTAGCAGTTGTTCTCCCCAAGTCAGAGCTCACTTGCCTTGAC  
451 S A L I D G S Q S T D D T E I V S Y H W  
1861 GTCAGCCCTCATTGATGGCAGCAAAGTACAGATGATACTGAAATAGTGAGTTATCATTG  
471 E E I N G P F I E E K T S V D S P V L R  
1921 GGAAGAAATAAACGGGCCCTCATAGAAGAGAAGACTTCAGTTGACTCTCCGTCTACG  
491 L S N L D P G N Y S F R L T V T D S D G  
1981 CTTGTCTAACCTTGATCCTGGTAACTATAGTTCAGGTTGACTGTTACAGACTCGGACGG  
511 A T N S T T A A L I V N N A V D Y P P V  
2041 AGCCACTAACTCTACAACTGCAGCCCTAATAGTGAACAATGCTGTGGACTACCCACCAAGT  
531 A N A G P N H T I T L P Q N S I T L N G  
2101 TGCTAATGCAGGACCAAATCACACCATAACTTGCCTCCAAACTCCATCACTTGAATGG  
551 N Q S S D D H Q I V L Y E W S L G P G S  
2161 AAACCAGAGCAGTGACGATCACCAGATTGTCTCTATGAGTGGTCCCTGGTCCTGGGAG  
571 E G K H V V M Q G V Q T P Y L H L S A M  
2221 TGAGGGCAAACATGTGGTCATGCAGGGAGTACAGACGCCATACCTTCAATTATCTGCAAT  
591 Q E G D Y T F Q L K V T D S S R Q Q S T  
2281 GCAGGAAGGAGATTATACATTCAGCTGAAGGTGACAGATCTTCAAGGCAACAGTCTAC  
611 A V V T V I V Q P E N N R P P V A V A G  
2341 TGCTGTAGTGACTGTGATTGTCCAGCCTGAAAACAATAGACCTCCAGTGGCTGTGGCCGG  
631 P D K E L I F P V E S A T L D G S S S S  
2401 CCCTGATAAAAGAGCTGATCTTCCCAGTGGAAAGTGCTACCCCTGGATGGAGCAGCAGCAG  
651 D D H G I V F Y H W E H V R G P S A V E

2461 CGATGACCACGGCATTGTCTTCTACCACTGGGAGCACGTCAAGAGGCCAGTGCAGTGG  
671 M E N I D K A I A T V T G L Q V G T Y H  
2521 GATGGAAAATATTGACAAAGCAATAGCCACTGTGACTGGTCTCCAGGTGGGACCTACCA  
691 F R L T V K D Q Q G L S S T S T L T V A  
2581 CTTCCGTTGACAGTGAAGAACAGCAGCAGGGACTGAGCAGCACGTCCACCCCTCACTGTGGC  
711 V K K E N N S P P R A R A G G G R H V L V  
2641 TGTGAAGAAGGAAAATAATAGTCCTCCCAGAGCCCAGGCTGGTGGCAGACATGTTCTTGT  
731 L P N N S I T L D G S R S T D D Q R I V  
2701 GCTTCCAATAATTCCATTACTTGGATGGTCAAGGTCTACTGATGACCAAAAGAATTGT  
751 S Y L W I R D G Q S P A A G D V I 'D G S  
2761 GTCCTATCTGTGGATCCGGATGGCCAGAGTCCAGCAGCTGGAGATGTCATCGATGGCTC  
771 D H S V A L Q L T N L V E G V Y T F H L  
2821 TGACCACAGTGTGGCTCTGCAGCTACGAATCTGGTGGAGGGGGTGTACACTTCCACTT  
791 R V T D S Q G A S D T D T A T V E V Q P  
2881 GCGAGTCACCGACAGTCAGGGGCCCTCGGACACAGACACTGCCACTGTGGAAGTGCAGCC  
811 D P R K S G L V E L T L Q V G V G Q L T  
2941 AGACCCCTAGGAAGAGTGGCTGGAGCTGACCTGCAAGGTTGGTGGCAGCTGAC  
831 E Q R K D T L V R Q L A V L L N V L D S  
3001 AGAGCAGCGGAAGGACACCTTGTGAGGCAGCTGGCTGTGCTGCTGAACGTGCTGGACTC  
851 D I K V Q K I R A H S D L S T V I V F Y  
3061 GGACATTAAGGTCCAGAAGATTGGGCCACTCGGATCTCAGCACCGTATTGTGTTTA  
871 V Q S R P P F K V L K A A E V A R N L H  
3121 TGTACAGAGCAGGCCCTTCAAGGTCTCAAAGCTGCTGAAGTGGCCGAAATCTGCA  
891 M R L S K E K A D F L L F K V L R V D T  
3181 CATGCGGCTCTCAAAGGAGAAGGCTGACTTCTGCTTCAAGGTCTTGAGGGTTGATAC  
911 A G C L L K C S G H G H C D P L T K R C  
3241 AGCAGGTTGCCTCTGAAGTGTCTGGCCATGGTCACTGGCACCCCTCACAAAGCGCTG  
931 I C S H L W M E N L I Q R Y I W D G E S  
3301 CATTGCTCTCACTTATGGATGGAGAACCTTATACAGCGTTATCTGGATGGAGAGAG  
951 N C E W S I F Y V T V L A F T L I V L T  
3361 CAACTGTGAGTGGAGTATATTCTATGTGACAGTGTGGCTTTACTCTTATTGTGCTAAC  
971 G G F T W L C I C C C K R Q K R T K I R  
3421 AGGAGGTTCACTTGGCTTGATCTGCTGCTGCAAAAGACAAAAAGGACTAAATCAG  
991 K K T K Y T I L D N M D E Q E R M E L R  
3481 GAAAAAAACAAAGTACACCATCCTGGATAACATGGATGAACAGGAAGAATGGAACGTGAG  
1011 P K Y G I K H R S T E H N S S L M V S E  
3541 GCCCAAATATGGTATCAAGCACCGAAGCACAGAGCACAACTCCAGCCTGATGGTATCCGA  
1031 S E F D S D Q D T I F S R E K M E R G N  
3601 GTCTGAGTTGACAGTGACAGGACACAATCTTCAGCCAGAGAAAGATGGAGAGAGGGAA  
1051 P K V S M N G S I R N G A S F S Y C S K  
3661 TCCAAAGGTTCCATGAATGGTCCATCAGAAATGGAGCTCCTTCAGTTATTGCTCAA  
1071 D R \*

3721 GGACAGATAATggcgcagttcattgtaaagtggaaaggacccttgaatccaagaccagtc  
3781 agtgggagttacagcacaaaacccactctttagaatagttcattgaccccttccccag  
3841 tgggttagatgttatccccacgtactaaaagaccgggtttgaaggcacaaaacaaaa  
3901 ct当地ttaactgagatgctgttaatagaaataaaggctggtaaaactctaagg  
3961 tatatactaaaagagtttgagttttagctggcacaatctcatattaaagatgaac  
4021 aacgatttctatctgttagaaccttagagaaggtgaatgaaacaaggttaaaaggat  
4081 gatttctgtcttagccgctgtgattgcctctaaggaacagcattctaaacacgggttctc  
4141 ttgttaggacactgcagtcagatggctgttatgttaaaatagcttgcattgtcaagaggcacgg  
4201 ccatctgtggaggtacggagtctgcattgttagcaagcttctgtgctgacggcaacactc  
4261 gcacagtgccaagccctcctgggttttaattctgtcattgtcaatggcagtttcatct  
4321 ctctcaagaaagcagctgtggcattcaagagctaaggagaatcgtattctaaggact  
4381 gaggcaatagaaaggggaggaggttaatgccgtcaggttgaaggtagcattgtaac  
4441 attatctttcttcttaagaaaaactacactgactcctctcggtgttagcagta  
4501 tagttctctaattgtaaacggatccccagttacattaaatgcaatagaagtgattaattc  
4561 attaagcatttattatgttctgttaggctgtcggttgactgccatagataggataacg  
4621 actcagcaattgttatattccaaaactctgaaatacagtcttgcattgttggatgg  
4681 cgtggttatgataactctggtccccgacaggtaacttccaaaataacttgcacatagatgt  
4741 ttcacttcatatgtttaaaaatcacatttaagtttctaccgaataatcttatttcaaa  
4801 catgaaagacaattaaaacattcccacccacaaagcagtaactcccgagcaattaactgg  
4861 gtttaatttgcctgtacgttactgggtcaggtagttcccatccacccttggcct  
4921 gaggctggtggccttgggtggcccttggcatttttggaaagattagaatgagagat  
4981 agaaccagtgtgtggtaccaagtgtgagcacacctaacaatatcctgtgcacaatgc  
5041 ttttttaacacatggaaaacttaggaatgcattgtcatgttgcatttttttttt  
5101 accagggcaggagtgcacagaaaaatgtttcccatgggttcttaaaaaaaaaattcagct  
5161 ttaggtgctttgtcatctcccgagtttgcatttgcatttttttttttttttttt  
5221 attgttaatttactgggaaaggcagaactaaaaatgtgtcattttttttttttttt  
5281 tgcttgcttatgcctacacttttgcatttgcatttgcatttttttttttttttttt  
5341 agaaggaaaatgtgattttttttttttttttttttttttttttttttttttttt  
5401 ccttatcaggtgaccagggttatggttttgcatttttttttttttttttttttttt  
5461 gacagcagccaaatgtaaagtcatggcgtaatgagggaaaggaggttgcatttt  
5521 cgctttatgtacataacatatgcatttttttttttttttttttttttttttttt  
5581 aagaggagatactatcatttttttttttttttttttttttttttttttttttttt  
5641 gtaacttcccgagggtcatggccaaaggactgttttttttttttttttttttttt  
5701 taaaactgttagttctgtttccactaaacaaaaggactcctgccttgcatttttttt  
5761 aaattctggtggaaactttttggccacctgaaagtcttgcatttttttttttttt  
5821 cttttatggatccagagagccaaagggtcagaggagagatggcctgcatttttttt  
5881 gatcacacccggccacccctcccttaggtttacagtggacttcttgccttgcatttt  
5941 ttttctgtccttggccatctcagccctggctctgtatccttgcatttttttttttt  
6001 gaatctctggaaatcaaacatcacagtagtgcatttttttttttttttttttt  
6061 ccatttctcatcagaacaaaagcagcagatggaaatgaccaaccagcatttttt  
6121 ctgcttatcattgaggatcttggagataaagcagcatttttttttttttttttt  
6181 acaggccctagaatatgggagtggttttttttttttttttttttttttttttttt

6241 ttgctggttacattcaatgaaggaggattcataccatggcattacaaggctaaacatg  
6301 tgtatgactaaggaactatctgaaaaacatgcagcaggtaagaaaatgtaccactcaac  
6361 aagccagtgtatgccaccctttgtgcgcggggaggagagtactaccattgttttgc  
6421 gacaaagctatcatggactatttatcttgggtttattgtctaaaatataatttttc  
6481 cctatgtgttgcataaggatattctaatatcacactattaaatataatgcactaatctaaat  
6541 aaagggtgtatgtatattctgtatgcatttttagggggaaatttgccttgc  
6601 tcagggttagaggattcccttgagtataggtcagcaactctggcctgcagcctgtgt  
6661 gcacgccccatgagccaaaatgtggcttatgtttcaatggtaaaaataaaaaaaa  
6721 aaatttgcataatgtgaactatatacattcagattgtgttcataaataaaatgttatt  
6781 ggaacatatacc

**Figure 2C. The cDNA (SEQ ID NO. : 6) and amino acid sequence (SEQ ID NO. : 7) of 254P1D6B v.3. The start methionine is underlined. The open reading frame extends from nucleic acid 739-3930 including the stop codon.**

1 gctgccgcggcggtggcgggatcccccggggtcaacccgtccacctgtgtgc  
61 cctccggcgggcctggctggcccgccgcagagccgcggcgctgcgtcactgcccga  
121 ggtgagagcgcagcagttagtccgcgttgcgttgcgttgcgttgcgttgc  
181 ggctacgtcccgaaaaagagggaaagagggaaagcggaggatttgcgtgggtgggtgttgc  
241 agcagggtgcgcgcgcgagggtgtgaacgtgtgtgtgtgtgtgtgtgtgt  
301 taagacctgcgtgcgtgcgcgcgcgcgcgcgcgcgcgcgcgcgcgcgc  
361 cagcaacgc  
421 tcccaagc  
481 acaaaaatgtggatgcgcgcgcgcgcgcgcgcgcgcgcgcgcgcgc  
541 aaaaacatcaaaaggaaagcttgcaccaactcttcaggccgcctcagaagcctgc  
601 caccactgtgtgtgcacaatggccgcgcgcgcgcgcgcgcgcgcgc  
661 tggtgacaatttcgcgtttgcgttgcactcatggcaaaaaatcactggtgag  
1 M T R L G W P S P C C A R K  
721 catcattaagaagacccATGACTAGACTGGCTGGCCGAGCCATGTTGTGCCGTAAAG  
15 Q C S E G R T Y S N A V I S P N L E T T  
781 CAGTGCAGCGAGGGAGGGACATATTCAATGCAGTCATTCACTAACGGAAACCA  
35 R I M R V S H T F P V V D C T A A C C D  
841 AGAACATGCAGGGGTCTCACACCTCCCTGCGTAGACTGCACGGCCGCTGCTGTGAC  
55 L S S C D L A W W F E G R C Y L V S C P  
901 CTGTCCAGCTGTGACCTGGCTGGTGGTCGAGGGCCGCTGCTACCTGGTGAGCTGCC  
75 H K E N C E P K K M G P I R S Y L T F V  
961 CACAAAGAGAACTGTGAGGCCAAGAAGATGGGCCCATCAGGTCTTATCTCACTTTGTG  
95 L R P V Q R P A Q L L D Y G D M M L N R  
1021 CTCCGGCCTGTTCAGAGGCCCTGCACAGCTGCTGGACTATGGGACATGATGCTAACAGG  
115 G S P S G I W G D S P E D I R K D L P F  
1081 GGCTCCCCCTCGGGGATCTGGGGGACTCACCTGAGGATATCAGAAAGGACTTGCCCTT  
135 L G K D W G L E E M S E Y S D D Y R E L  
1141 CTAGGCAAAGATTGGGCCTAGAGGAGATGTCTGAGTACTCAGATGACTACCGGGAGCTG

155 E K D L L Q P S G K Q E P R G S A E Y T  
1201 GAGAAGGACCTCTTGCAACCCAGTGGCAAGCAGGAGGCCAGAGGGAGTGCCGAGTACACG  
175 D W G L L P G S E G A F N S S V G D S P  
1261 GACTGGGGCCTACTGCCGGCAGCAGGGGGCCTCAACTCCTCTGTTGGAGACAGTCCT  
195 A V P A E T Q Q D P E L H Y L N E S A S  
1321 GCGGTGCCAGCGGAGACGCAGCAGGACCTGAGCTCCATTACCTGAATGAGTCGGCTTCA  
215 T P A P K L P E R S V L L P L P T T P S  
1381 ACCCCTGCCCAAAACTCCCTGAGAGAAGTGTGTTGCTTCCCTGCCGACTACTCCATCT  
235 S G E V L E K E K A S Q L Q E Q S S N S  
1441 TCAGGAGAGGTGTTGGAGAAAGAAAAGGCTCTCAGCTCCAGGAACAATCCAGCAACAGC  
255 S G K E V L M P S H S L P P A S L E L S  
1501 TCTGGAAAAGAGGTTCTAATGCCCTCCCAGTCAGCTCCGGCAAGCCTGGAGCTCAGC  
275 S V T V E K S P V L T V T P G S T E H S  
1561 TCAGTCACCGTGGAGAAAAGCCCAGTGCTCACAGTCACCCGGGAGTACAGAGCACAGC  
295 I P T P P T S A A P S E S T P S E L P I  
1621 ATCCCAACACCTCCCACTAGCGCAGCCCCCTCTGAGTCCACCCCATCTGAGCTACCCATA  
315 S P T T A P R T V K E L T V S A G D N L  
1681 TCTCCTACCCTGCTCCAGGACAGTGAAAGAACTTACGGTATCGGCTGGAGATAACCTA  
335 I I T L P D N E V E L K A F V A P A P P  
1741 ATTATAACTTACCCGACAATGAAGTTGAACTGAAGGCCTTGTGCGCCAGCGCCACCT  
355 V E T T Y N Y E W N L I S H P T D Y Q G  
1801 GTAGAAACAACCTACAACATGAATGGAATTAAAGCCACCCACAGACTACCAAGGT  
375 E I K Q G H K Q T L N L S Q L S V G L Y  
1861 GAAATAAAACAAGGACACAAGCAAACCTTAACCTCTCAATTGTCCGTCGGACTTTAT  
395 V F K V T V S S E N A F G E G F V N V T  
1921 GTCTTCAAAGTCACTGTTCTAGTGAAACGCCCTTGGAGAAGGATTGTCAATGTCACT  
415 V K P A R R V N L P P V A V V S P Q L Q  
1981 GTTAAGCCTGCCAGAAGAGTCACCTGCCACCTGTAGCAGTTGTTCTCCCCAACTGCAA  
435 E L T L P L T S A L I D G S Q S T D D T  
2041 GAGCTCACTTGCCTTGACGTCAAGCCCTCATTGATGGCAGCCAAAGTACAGATGATACT  
455 E I V S Y H W E E I N G P F I E E K T S  
2101 GAAATAGTGAGTTATCATGGGAAGAAATAACGGGCCCTCATAGAAGAGAAAGACTTCA  
475 V D S P V L R L S N L D P G N Y S F R L  
2161 GTTGACTCTCCGTCTACGCTTGATCCTGGTAACTATAGTTCAGGTTG  
495 T V T D S D G A T N S T T A A L I V N N  
2221 ACTGTTACAGACTCGGACGGAGCCACTAACTCTACAACTGCAGCCCTAATAGTGAACAAT  
515 A V D Y P P V A N A G P N H T I T L P Q  
2281 GCTGTGGACTACCCACCAGTTGCTAATGCAGGACCAAATCACACCATAACTTGCCCCAA  
535 N S I T L N G N Q S S D D H Q I V L Y E  
2341 AACTCCATCACTTGAATGGAAACCAGAGCAGTGACGATCACCAGATTGTCCCTATGAG  
555 W S L G P G S E G K H V V M Q G V Q T P  
2401 TGGTCCCTGGGTCTGGAGTGAGGGCAAACATGTGGTCATGCAGGGAGTACAGACGCCA

575 Y L H L S A M Q E G D Y T F Q L K V T D  
2461 TACCTTCATTATCTGCAATGCAGGAAGGAGATTATACATTTCAAGCTGAAGGTGACAGAT  
595 S S R Q Q S T A V V T V I V Q P E N N R  
2521 TCTTCAAGGCAACAGTCACTGCTGGTGACTGTGATTGTCCAGCCTGAAAACAATAGA  
615 P P V A V A G P D K E L I F P V E S A T  
2581 CCTCCAGTGGCTGTGGCCGGCCCTGATAAAAGAGCTGATCTTCCCAGTGGAAAGTGCTACC  
635 L D G S S S S D D H G I V F Y H W E H V  
2641 CTGGATGGGAGCAGCAGCGATGACCACGGCATTGTCTTCTACCACTGGGAGCAGTC  
655 R G P S A V E M E N I D K A I A T V T G  
2701 AGAGGCCCAAGTGCAGTGGAGATGAAAATATTGACAAAGCAATAGCCACTGTGACTGGT  
675 L Q V G T Y H F R L T V K D Q Q G L S S  
2761 CTCCAGGTGGGGACCTACCACCTCGTTGACAGTGAAAGACCAGCAGGACTGAGCAGC  
695 T S T L T V A V K K E N N N S P P R A R A  
2821 ACGTCCACCCCACTGTGGCTGTGAAGAAGGAAAATAATAGTCCTCCCAGAGCCGGCT  
715 G G R H V L V L P N N S I T L D G S R S  
2881 GGTGGCAGACATGTTCTGTGCTTCCAATAATTCCATTACTTTGGATGGTCAAGGTCT  
735 T D D Q R I V S Y L W I R D G Q S P A A  
2941 ACTGATGACCAAAGAATTGTGCTTATCTGTGGATCCGGATGGCAGACTCCAGCAGCT  
755 G D V I D G S D H S V A L Q L T N L V E  
3001 GGAGATGTCATCGATGGCTCTGACCACAGTGTGGCTCTGCAGCTTACGAATCTGGTGGAG  
775 G V Y T F H L R V T D S Q G A S D T D T  
3061 GGGGTGTACACTTCCACTTGCAGTCACCGACAGTCAGGGGGCCTCGGACACAGACACT  
795 A T V E V Q P D P R K S G L V E L T L Q  
3121 GCCACTGTGAAAGTGCAGCCAGACCCCTAGGAAGAGTGGCTGGAGCTGACCCCTGCAG  
815 V G V G Q L T E Q R K D T L V R Q L A V  
3181 GTTGGTGTGAGCTGACAGAGCAGCGGAAGGACACCCCTGTGAGGCAGCTGGCTGTG  
835 L L N V L D S D I K V Q K I R A H S D L  
3241 CTGCTGAACGTGCTGGACTCGGACATTAAGGTCCAGAAGATTGGGCCACTCGGATCTC  
855 S T V I V F Y V Q S R P P F K V L K A A  
3301 AGCACCGTATTGTGTTATGTACAGAGCAGGCCCTTCAAGGTTCTCAAAGCTGCT  
875 E V A R N L H M R L S K E K A D F L L F  
3361 GAAGTGGCCGAAATCTGCACATGGCTCTCAAAGGAGAAGGCTGACTTCTGCTTTC  
895 K V L R V D T A G C L L K C S G H G H C  
3421 AAGGTCTTGAGGGTTGATAACAGCAGGTTGCCTCTGAAGTGTCTGGCATGGTCACTGC  
915 D P L T K R C I C S H L W M E N L I Q R  
3481 GACCCCTCACAAAGCGCTGCATTGCTCTCAATTGATGGAGAACCTTATAACAGCGT  
935 Y I W D G E S N C E W S I F Y V T V L A  
3541 TATATCTGGATGGAGAGAGCAACTGTGAGTGGAGTATATTCTATGTGACAGTGGCT  
955 F T L I V L T G G F T W L C I C C C K R  
3601 TTTACTCTTATTGTGCTAACAGGAGGTTCACTTGGCTTGCATCTGCTGCTGCAAAAGA  
975 Q K R T K I R K K T K Y T I L D N M D E  
3661 CAAAAAAGGACTAAAATCAGGAAAAAAACAAAGTACACCCTGGATAACATGGATGAA

995 Q E R M E L R P K Y G I K H R S T E H N  
3721 CAGGAAAGAACGGAACTGAGGCCAAATATGGTATCAAGCACCAGCACAGAGCACAAAC  
1015 S S L M V S E S E F D S D Q D T I F S R  
3781 TCCAGCCTGATGGTATCCGAGTCTGAGTTGACAGTGACCGAGCACAAATCTCAGCCGA  
1035 E K M E R G N P K V S M N G S I R N G A  
3841 GAAAAGATGGAGAGAGGGAATCCAAAGGTTCCATGAATGGTTCCATCAGAAATGGAGCT  
1055 S F S Y C S K D R \*  
3901 TCCTTCAGTTATTGCTCAAAGGACAGATAAtggcgcaagttcattgtaaagtggaggacc  
3961 ccttgaatccaagaccagtcaatggggatgttacagcacaaaacccactcttttagaatagt  
4021 tcattgaccttcttcccagtgggttagatgttatcccacgtactaaaagaccgggtt  
4081 ttgaaggcacaaaaacaaaactttgtcttttaactgagatgctgttaatagaaaataaa  
4141 ggctggtaaaaactctaaggatataactttaaaagagttttgagtttttagctggcaca  
4201 atctcatattaaagatgaacaacgattctatctgttagaaccccttagagaaggtaatgaa  
4261 acaaggtttaaaaaggatgatttctgtcttagccgctgtgattgcctctaaggAACAG  
4321 cattctaaacacgggttctgttaggacctgcagtcagatggctgttatgttaaaaata  
4381 gcttgtctaagaggcacggccatctgtggaggtacggagtctgcattgtcaagcttt  
4441 ctgtgctgacggcaacactcgcacagtgcacagccctctgggttttaattctgtctat  
4501 gtcaatggcagtttcatctctcaagaaagcagctgtggccattcaagagactaaggaa  
4561 agaatcgattctaaggactgaggcaatagaaaaggggaggaggacttaatgcgtgcag  
4621 gttgaaggtagcattgttaacattatctttcttcttaagaaaaactacactgactcct  
4681 ctcgggttttagcagttatgttcttaatgttaacccatgttttttttttttttttttt  
4741 gcaatagaagtgattaattcattaaggactcatttttttttttttttttttttttttt  
4801 tgccatagataggataacgactcagcaattgttatattccaaaactctgaaataaca  
4861 gtcagtttactggatggcgtggatgtataactctggccccgacaggactttccaa  
4921 aataacttgacatagatgttattcatttttttttttttttttttttttttttttttt  
4981 cgaataaatcttatttcaaacatgaaaagacaatttttttttttttttttttttttt  
5041 ctcccgagcaatt  
5101 ccccatccacccttggcttgaggctggcgtggcatttttttttttttttttttttt  
5161 ggaagatttagaatgagagatagaaccaggactgttttttttttttttttttttttt  
5221 aatatt  
5281 aagaagcaaggatt  
5341 tcttt  
5401 ggaccatcttatt  
5461 catt  
5521 tcaataactgtctatgttttttttttttttttttttttttttttttttttttttt  
5581 cttcataaggctagaatctgccttattcaggtgaccagggttatgggttttgcatt  
5641 tggatatt  
5701 agaaggagtgaaacatttaccgctttatgtacataacatgtcagtttacataact  
5761 gatccttataatcaaccccttgaagaggagatactatcatttttttttttttttt  
5821 tgaaggcccagagaggtaagtaacttccagaggcatggccaagaaggtagtggtt  
5881 agaactgaatgcaaatt  
5941 tgccttgcatt

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6001 cccaggactaagaggaatttttaatggatccagagagccaaggcagaggagat
6061 ggcctgcatagtctcctgtggatcacaccggccaccctcccttagttacagtgg
6121 acttcttctgcccctccctttctgtccttgcacatctcagcctggctctgtatcc
6181 ttccatcacagaaggatcttgcataatctctggaaatcaaacatcacagtagtgcata
6241 gtgagtcctgtcttgcacccattctcatcagaacaaggcacgagatggaaatgacca
6301 ccagcattcttcatggtgactgcttacatttgcacatggaggatcttggagataaagcacgcta
6361 agagctctggacagagaaaaacaggccctagaatatggagtggtgttgcacggctca
6421 taggctaacaaggacttttagttgcgtgttacattcaatgcacggaggattcataccatg
6481 gcattacaaggctaaggcatgtgtatgactaaggaactatctgaaaacatgcacggat
6541 aagaaaatgtaccactcaacaaggccactgtatggttgcgcggggaggagatg
6601 actaccattgtttttgtgtgacaaagctatcatggactatttatcttgcgtttatttgc
6661 cttaaaatattatccatgtgtgacaaaggatgttgcataatatcacactattaa
6721 atatatgcactaatctaaataaagggtgttgcataatgtttgcgttgcataatgttttaggggg
6781 aaatttgcgtttatgtttgcgttgcacgcggatcccttgcgttgcataatgttttaggggg
6841 ctggcctgcagcctgtgtgcacgcggatcccttgcgttgcataatgttttaggggg
6901 atggtaaaaataaataaaaaatttgcataatgtgttgcgttgcataatgttttaggggg
6961 ttcataaataaaggatgtttatgtgttgcgttgcataatgttttaggggg

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**Figure 2D. 254P1D6B v.4 through v.20, SNP variants of 254P1D6B v.1.** The 254P1D6B v.4 through v.20 proteins have 1072 amino acids. Variants 254P1D6B v.4 through v.20 are variants with single nucleotide difference from 254P1D6B v.1. 254P1D6B v.5 and v.6 proteins differ from 254P1D6B v.1 by one amino acid. 254P1D6B v.4 and v.7 through v.20 proteins code for the same protein as v.1. Though these SNP variants are shown separately, they can also occur in any combinations and in any of the transcript variants listed above in Figures 2A, Figure 2B and Figure 2C.

Variant	Nucleic acid position	Nucleic Acid Variation	Amino Acid Position	Amino Acid Variation
254P1D6B v.4	286	C/G	Silent variant	
254P1D6B v.5	935	C/A	142	P=>T
254P1D6B v.6 (Identical AA as v.2)	980	T/G	157	S=>A
254P1D6B v.7	2347	G/A	Silent variant	
254P1D6B v.8	3762	C/T	Silent variant	
254P1D6B v.9	3772	A/G	Silent variant	
254P1D6B v.10	3955	C/T	Silent variant	
254P1D6B v.11	4096	C/T	Silent variant	
254P1D6B v.12	4415	G/A	Silent variant	
254P1D6B v.13	4519	G/A	Silent variant	
254P1D6B v.14	4539	A/G	Silent variant	

Variant	Nucleic Acid Position	Nucleic Acid Variation	Amino Acid Position	Amino Acid Variation
254P1D6B v.15	4614	G/T	<i>Silent variant</i>	
254P1D6B v.16	5184	G/C	<i>Silent variant</i>	
254P1D6B v.17	5528	T/G	<i>Silent variant</i>	
254P1D6B v.18	5641	G/A	<i>Silent variant</i>	
254P1D6B v.19	6221	T/C	<i>Silent variant</i>	
254P1D6B v.20	6223	G/A	<i>Silent variant</i>	

**Figure 3:**

**Figure 3A. Amino acid sequence 254P1D6B v.1 clone LCP-3 (SEQ ID NO.: 8).** The 254P1D6B v.1 clone LCP-3 protein has 1072 amino acids.

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1 MAPPTGVLS LLLLVTIAGC ARKQCSEGRT YSNAVISPRL ETTRIMRVSH TFPVVDCTAA
61 CCDLSSCDLA WWFEGRCYLV SCPHKENCEP KKMGPRLSYL TFVLRPVQRP AQLLDYGDMM
121 LNRGSPSGIW GDSPEDIRKD LPFLGKDGL EEMSEYADDY RELEKDLLQP SGKQEPRGSA
181 EYTDWGLLPG SEGAFNSSVG DSPAVPAETQ QDPELHYLNE SASTPAPKLP ERSVLLPLPT
241 TPSSGEVLEK EKASQLQEWS SNSSGKEVLM PSHSLPPASL ELSSVTVEKS PVLTVTPGST
301 EHSIPTPPPTS AAPSESTPSE LPISPTTAPR TVKELTVSAG DNLIITLPDN EVELKAFVAP
361 APPVETTYNY EWNLISHPTD YQGEIKQGHK QTLNLSQLSV GLYVFKVTVS SENAFGEGFV
421 NVTVKPARRV NLPPVAVVSP QLQELTLPLT SALIDGSQST DDTEIVSYHW EEINGPFIIE
481 KTSVDPVLR LSNLDPGNYS FRLTVTDSDG ATNSTTAALI VNNAVDYPPV ANAGPNHTIT
541 LPQNSITLNG NQSSDDHQIV LYEWLSLPGS EGKHHVVMQGV QTPYLHLSAM QEGDYTFQLK
601 VTDSSRQQST AVVTIVQPE NNRPPVAVAG PDKELEIFPVE SATLDGSSSS DDHGIVFYHW
661 EHVRGSAVE MENIDKAIAT VTGLQVGTYH FRLTVKDQQG LSSTSTLTVA VKKENNSPPR
721 ARAGGRHVLV LPNNISITLDG SRSTDDQRIV SYLWIRDGQS PAAGDVIDGS DHSVALQLTN
781 LVEGVYTFHL RVTDSQGASD TDTATVEVQP DPRKSGLVEL TLQVGVGQLT EQRKDTLVRQ
841 LAVLLNVLDS DIKVQKIRAH SDLSTVIVFY VQSRPPFKVL KAAEVARNLH MRLSKEKADF
901 LLFKVLRVDT AGCLLKCNSGH GHCDPLTKRC ICSHLWMENL IQRYIWDGES NCEWSIFYVT
961 VLAFTLIVLT GGFTWLCICC CKRQKRTKIR KKTQYTILDN MDEQERMELR PKYGIKRST
1021 EHNSSLMVSE SEFDSDQDTI FSREKMERGN PKVSMNGSIR NGASFSSYCSK DR
```

**Figure 3B. Amino acid sequence 254P1D6B v.2 (SEQ ID NO.: 9).** The 254P1D6B v.2 protein has 1072 amino acids.

```
1 MAPPTGVLS LLLLVTIAGC ARKQCSEGRT YSNAVISPRL ETTRIMRVSH TFPVVDCTAA
61 CCDLSSCDLA WWFEGRCYLV SCPHKENCEP KKMGPRLSYL TFVLRPVQRP AQLLDYGDMM
121 LNRGSPSGIW GDSPEDIRKD LPFLGKDGL EEMSEYADDY RELEKDLLQP SGKQEPRGSA
181 EYTDWGLLPG SEGAFNSSVG DSPAVPAETQ QDPELHYLNE SASTPAPKLP ERSVLLPLPT
241 TPSSGEVLEK EKASQLQEWS SNSSGKEVLM PSHSLPPASL ELSSVTVEKS PVLTVTPGST
301 EHSIPTPPPTS AAPSESTPSE LPISPTTAPR TVKELTVSAG DNLIITLPDN EVELKAFVAP
361 APPVETTYNY EWNLISHPTD YQGEIKQGHK QTLNLSQLSV GLYVFKVTVS SENAFGEGFV
421 NVTVKPARRV NLPPVAVVSP QLQELTLPLT SALIDGSQST DDTEIVSYHW EEINGPFIIE
481 KTSVDPVLR LSNLDPGNYS FRLTVTDSDG ATNSTTAALI VNNAVDYPPV ANAGPNHTIT
541 LPQNSITLNG NQSSDDHQIV LYEWLSLPGS EGKHHVVMQGV QTPYLHLSAM QEGDYTFQLK
601 VTDSSRQQST AVVTIVQPE NNRPPVAVAG PDKELEIFPVE SATLDGSSSS DDHGIVFYHW
661 EHVRGSAVE MENIDKAIAT VTGLQVGTYH FRLTVKDQQG LSSTSTLTVA VKKENNSPPR
721 ARAGGRHVLV LPNNISITLDG SRSTDDQRIV SYLWIRDGQS PAAGDVIDGS DHSVALQLTN
781 LVEGVYTFHL RVTDSQGASD TDTATVEVQP DPRKSGLVEL TLQVGVGQLT EQRKDTLVRQ
841 LAVLLNVLDS DIKVQKIRAH SDLSTVIVFY VQSRPPFKVL KAAEVARNLH MRLSKEKADF
901 LLFKVLRVDT AGCLLKCNSGH GHCDPLTKRC ICSHLWMENL IQRYIWDGES NCEWSIFYVT
```

961 VLAFTLIVLT GGFTWLCICC CKRKRTKIR KKTKYTILDN MDEQERMELR PKYGIKHRST  
1021 EHNSSLMVSE SEFDSDQDTI FSREKMERGN PKVSMNGSIR NGASF SYCSK DR

Figure 3C. Amino acid sequence 254P1D6B v.3 (SEQ ID NO: 10). The 254P1D6B v.3 protein has 1063 amino acids.

1 MTRLGWPSPC CARKQCSEGR TYSNAVISPN LETTRIMRVS HTFPVVDCTA ACCDLSSCDL  
61 AWWFEGRCYL VSCPHKENCE PKKMGPIRSY LTFVLRPVQR PAQLLDYGDM MLNRGSPSGI  
121 WGDSPEDIRK DLPFLGKDWG LEEMSEYSDD YRELEKDLLQ PSGKQEPRGS AEYTDWGLLP  
181 GSEGAFNSSV GDSPA VPAET QQDPELHYLN ESASTPAPKL PERSVLLPLP TTPSSGEVLE  
241 KEKASQLQEQ SSNSSGKEVL MPSHSLPPAS LELSSVTVEK SPVLT VTPGS TEHSIPTPPT  
301 SAAPSESTPS ELPISPTTAP RTVKELTVSA GDNLII TLPD NEVELKAFVA PAPPVETTYN  
361 YEWNLISHPT DYQGEIKQGH KQLNLSQLS VGLYVFKVTV SSENAFGE GF VNVTVKPARR  
421 VNLPPVAVVS PQQLQELTLPL TSALIDGSQS TDDTEIVSYH WEEINGPFIE EKTSVDPV  
481 RLSNLDPGNY SFRLTVTDSD GATNSTTAAL IVNNNAVDYPP VANAGPNHTI TLPQNSITLN  
541 GNQSSDDHQI VLYEWSLGPG SEGKHVVMQG VQTPYLHLSA MQEGDYTFQL KVTDSSRQQS  
601 TAVVTVIVQP ENNRPPVAVA GPDKELIFPV ESATLDGSSS SDDHGIVFYH WEHVRGPAV  
661 EME NIDKAIA TVTGLQVGTY HFR LTVKDQQ GLSSTSTLTV AVKKENN SPP RARAGGRHVL  
721 VLPNNSITLD GSRSTDDQRI VSYLWIRDGQ SPAAGDVIDG SDHSVALQLT NLVEGVYTFH  
781 LRVTDSQGAS DTDTATVEVQ PDPRKSGLVE LTLQVGVQQL TEQRKDTLVR QLAVLLNVLD  
841 SDIKVQKIRA HSDLSTVIVF YVQSRPPFKV LKAAEVARNL HMRLSKEKAD FLLFKVLRVD  
901 TAGCLLKCSG HGHCDPLTKR CICSHLWMEN LIQRYIWDGE SNCEWSIFYV TVLAFTLIVL  
961 TGGFTWLCIC CCKRKRTKIR RKKT KYTILD NMDEQERMEL RPKYGIKRS TEHNSSLMVS  
1021 ESEFDSDQDT IFSREKMERG NPKVSMNGSIR RNGASF SYCS KDR

Figure 3D. Amino acid sequence 254P1D6B v.5 (SEQ ID NO: 11). The 254P1D6B v.5 protein has 1072 amino acids.

1 MAPPTGV LSS LLLLVTIAGC ARKQCSEGRT YSNAVISP NL ETTRIMRVSH TFPVVDCTAA  
61 C C D L S S C D L A W W F E G R C Y L V S C P H K E N C E P K K M G P I R S Y L T F V L R P V Q R P A Q L L D Y G D M M  
121 L N R G S P S G I W G D S P E D I R K D L T F L G K D W G L E E M S E Y S D D Y R E L E K D L L Q P S G K Q E P R G S A  
181 E Y T D W G L L P G S E G A F N S S V G D S P A V P A E T Q Q D P E L H Y L N E S A S T P A P K L P E R S V L L P L P  
241 T P S S G E V L E K E K A S Q L Q E Q S S N S S G K E V L M P S H S L P P A S L E L S S V T V E K S P V L T V T P G S T  
301 E H S I P T P P T S A A P S E S T P S E L P I S P T T A P R T V K E L T V S A G D N L I I T L P D N E V E L K A F V A P  
361 A P P V E T T Y N Y E W N L I S H P T D Y Q G E I K Q G H K Q T L N L S Q L S V G L Y V F K V T V S S E N A F G E G F V  
421 N V T V K P A R R V N L P P V A V V S P Q L Q E L T L P L T S A L I D G S Q S T D D T E I V S Y H W E E I N G P F I E E  
481 K T S V D S P V L R L S N L D P G N Y S F R L T V T D S D G A T N S T T A A L I V N N A V D Y P P V A N A G P N H T I T  
541 L P Q N S I T L N G N Q S S D D H Q I V L Y E W S L G P G S E G K H V V M Q G V Q T P Y L H L S A M Q E G D Y T F Q L K  
601 V T D S S R Q Q S T A V V T V I V Q P E N N R P P V A V A G P D K E L I F P V E S A T L D G S S S S D D H G I V F Y H W  
661 E H V R G P A S V E M E N I D K A I A T V T G L Q V G T Y H F R L T V K D Q Q G L S S T S T L T V A V K K E N N S P P R  
721 A R A G G R H V L V L P N N S I T L D G S R S T D D Q R I V S Y L W I R D G Q S P A A G D V I D G S D H S V A L Q L T N  
781 L V E G V Y T F H L R V T D S Q G A S D T D T A T V E V Q P D P R K S G L V E L T L Q V G V G Q L T E Q R K D T L V R Q  
841 L A V L L N V L D S D I K V Q K I R A H S D L S T V I V F Y V Q S R P P F K V L K A A E V A R N L H M R L S K E K A D F  
901 L L F K V L R V D T A G C L L K C S G H G H C D P L T K R C I C S H L W M E N L I Q R Y I W D G E S N C E W S I F Y V T

961 VLAFTLIVLT GGFTWLCICC CKRQKRTKIR KKTKYTILDN MDEQERMELR PKYGIKHRST  
1021 EHNSSLMVSE SEFDSDQDTI FSREKMERGN PKVSMNGSIR NGASF SYCSK DR

Figure 3E. Amino acid sequence 254P1D6B v.6 (SEQ ID NO: 12). The 254P1D6B v.6 protein has 1072 amino acids.

1 MAPPTGVLS LLLLVTIAGC ARKQCSEGRT YSNAVISPNL ETRTRIMRVSH TFPVVVDCTAA  
61 CCDLSSCDLA WWFEGRCYLV SCPHKENCEP KKMGPIRSYL TFVLRPVQRP AQLLDYGDMM  
121 LNRGSPSGIW GDSPEDIRKD LPFLGKDHWGL EEMSEYADDY RELEKDLLQP SGKQEPRGSA  
181 EYTDWGLLPG SEGAFNNSVG DSPAVPAETQ QDPELHYLNE SASTPAPKLP ERSVLLPLPT  
241 TPSSGEVLEK EKASQLQEQS SNSSGKEVLM PSHSLPPASL ELSSTVTEKS PVLTVTGOST  
301 EHSIPTPPPTS AAPSESTPSE LPISPTTAPR TVKELTVSAG DNLIITLPDN EVELKAFVAP  
361 APPVETTYNY EWNLISHPTD YQGEIKQGHK QTLNLSQLSV GLYVFKVTVS SENAFGEGFV  
421 NVTVKPARRV NLPPVAVVSP QLQELTLPLT SALIDGSQST DDTEIVSYHW EEINGPFIEE  
481 KTSVDSPVLR LSNLDPGNYS FRLTVTDSDG ATNSTTAALI VNNAVDYPVV ANAGPNHTIT  
541 LPQNSITLNG NQSSDDHQIV LYEWLSLPGS EGKHVVMQGV QTPYLHLSAM QEGDYTFQLK  
601 VTDSSRQQST AVVTVIVQPE NNRPPVAVAG PDKELIFPVE SATLDGSSSS DDHGIVFYHW  
661 EHVRGSAVE MENIDKAIAT VTGLQVGTYH FRLTVKDQQG LSSTSTLTVA VKKENNSPPR  
721 ARAGGRHV LV LPNNSITLDG SRSTDQDQIV SYLWIRDQGS PAAGDVIDGS DHSVALQLTN  
781 LVEGVYTFHL RVTDSQGASD TDTATVEVQP DPRKSGLVEL TLQVGVGQLT EQRKDTLVRQ  
841 LAVLLNVLDS DIKVQKIRAH SDLSTVIVFY VQSRPPFKVL KAAEVARNLH MRLSKEKADF  
901 LLFKVLRVDT AGCLLKCSCGH GHCDPLTKRC ICSHLWMENL IQRYIWDGES NCEWSIFYVT  
961 VLAFTLIVLT GGFTWLCICC CKRQKRTKIR KKTKYTILDN MDEQERMELR PKYGIKHRST  
1021 EHNSSLMVSE SEFDSDQDTI FSREKMERGN PKVSMNGSIR NGASF SYCSK DR

Figure 5: 254P1D6B variant 1

Hydrophilicity profile

(Hopp T.P., Woods K.R., 1981.

Proc. Natl. Acad. Sci. U.S.A. 78:3824-3828)

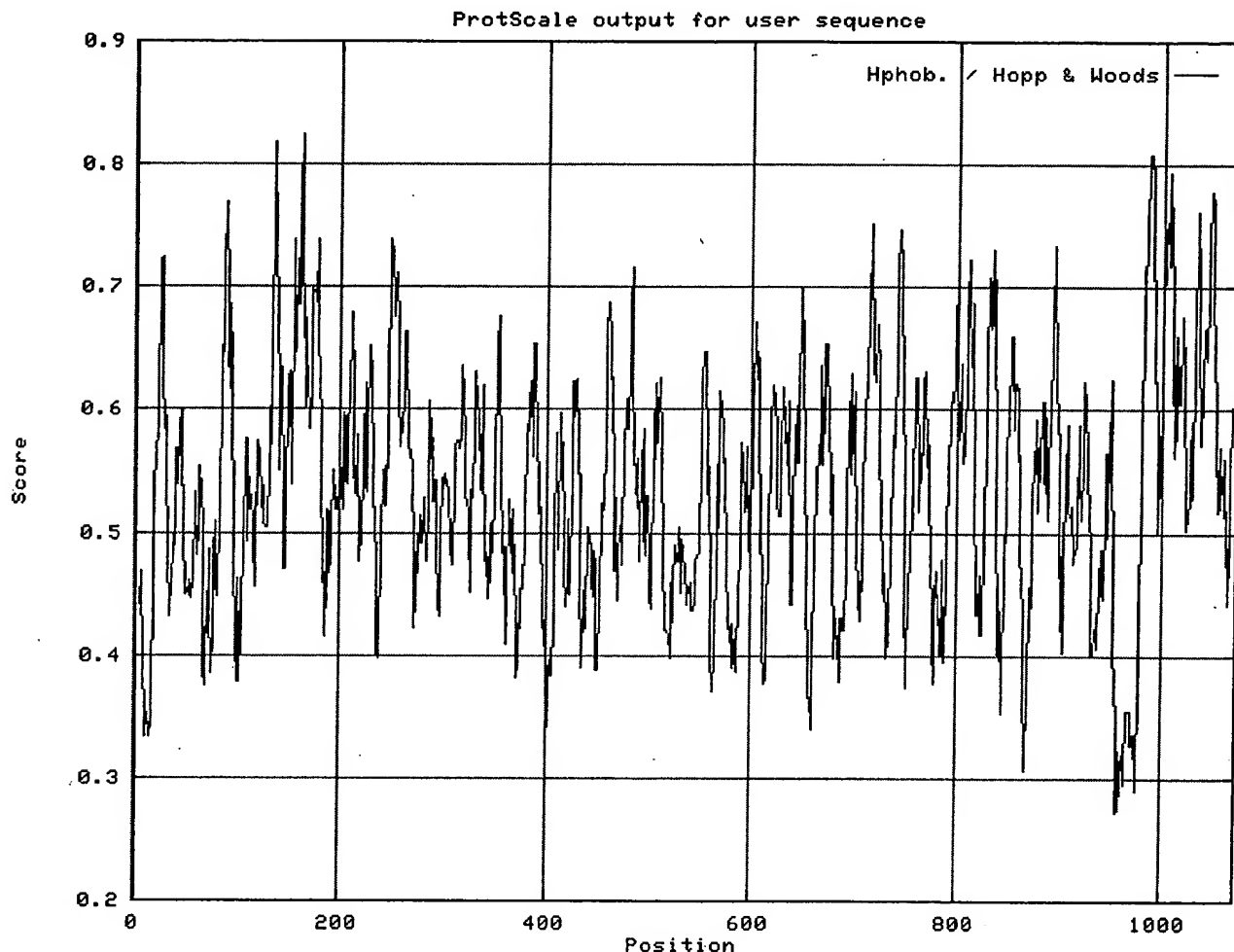
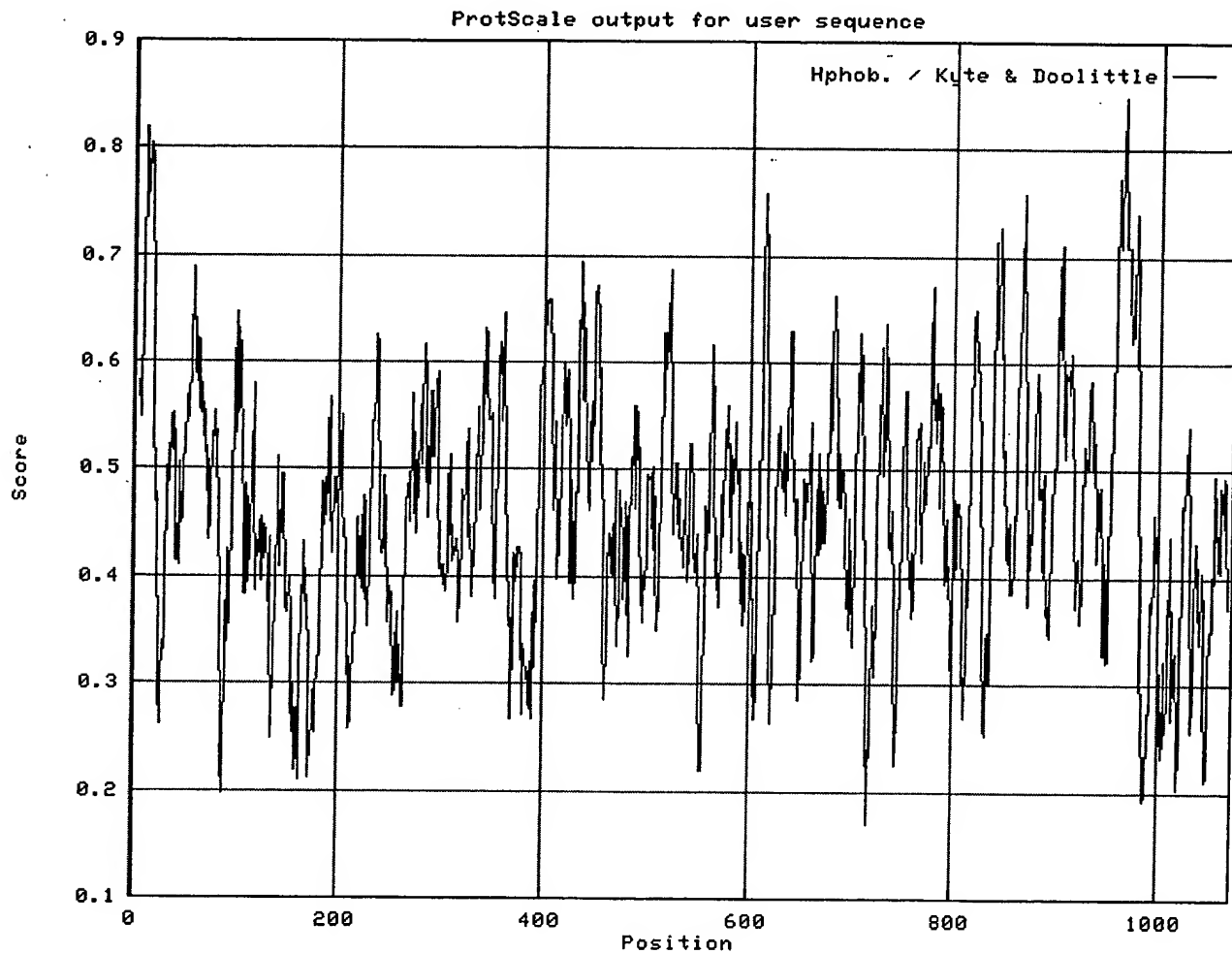
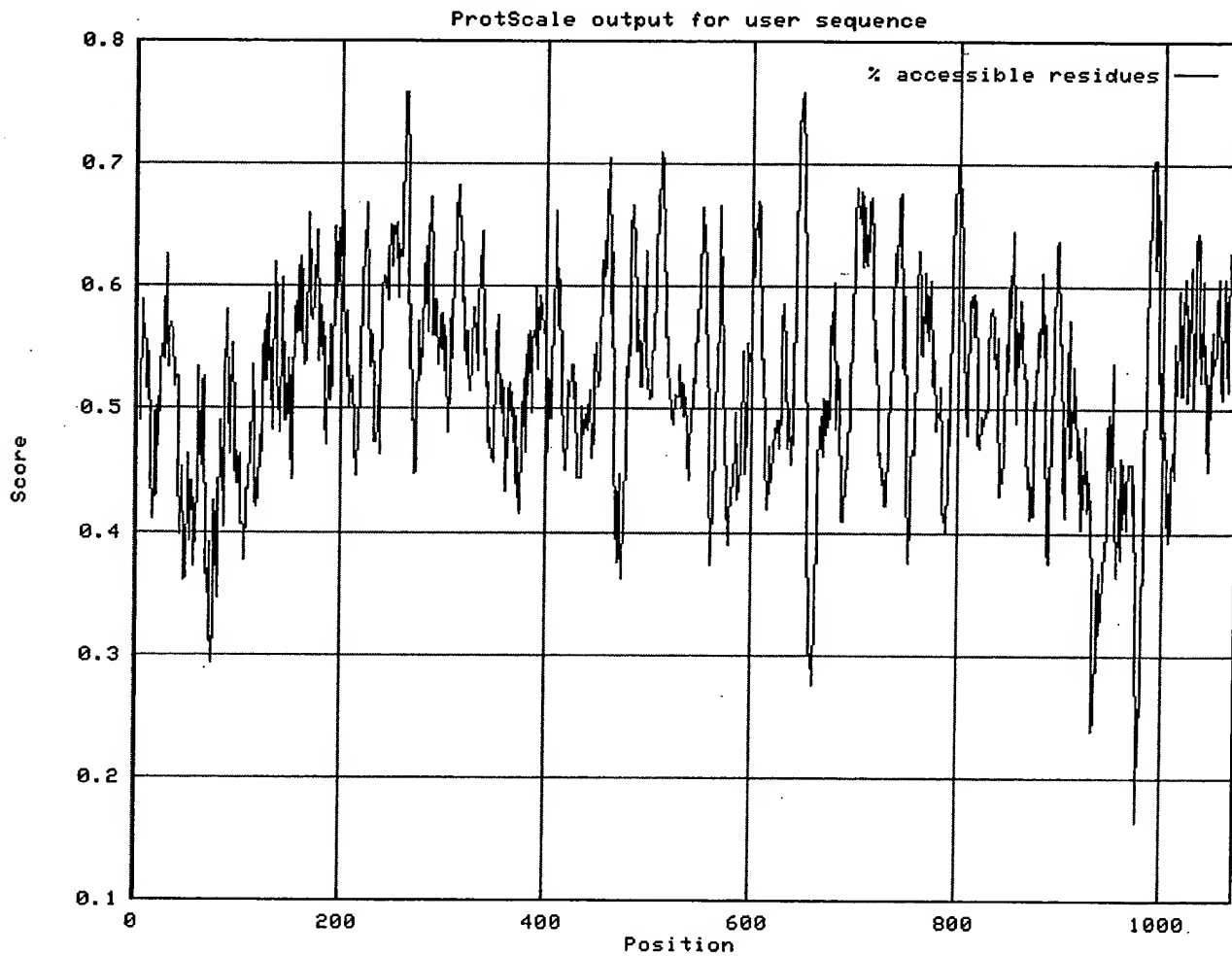


Figure 6: 254P1D6B variant 1  
Hydropathicity Profile  
(Kyte J., Doolittle R.F., 1982. J. Mol. Biol. 157:105-132)



# Figure 7: 254P1D6B variant 1 % Accessible Residues Profile (Janin J., 1979. Nature 277:491-492)



# Figure 8: 254P1D6B variant 1

## Average Flexibility Profile

(Bhaskaran R., Ponnuswamy P.K., 1988.  
Int. J. Pept. Protein Res. 32:242-255)

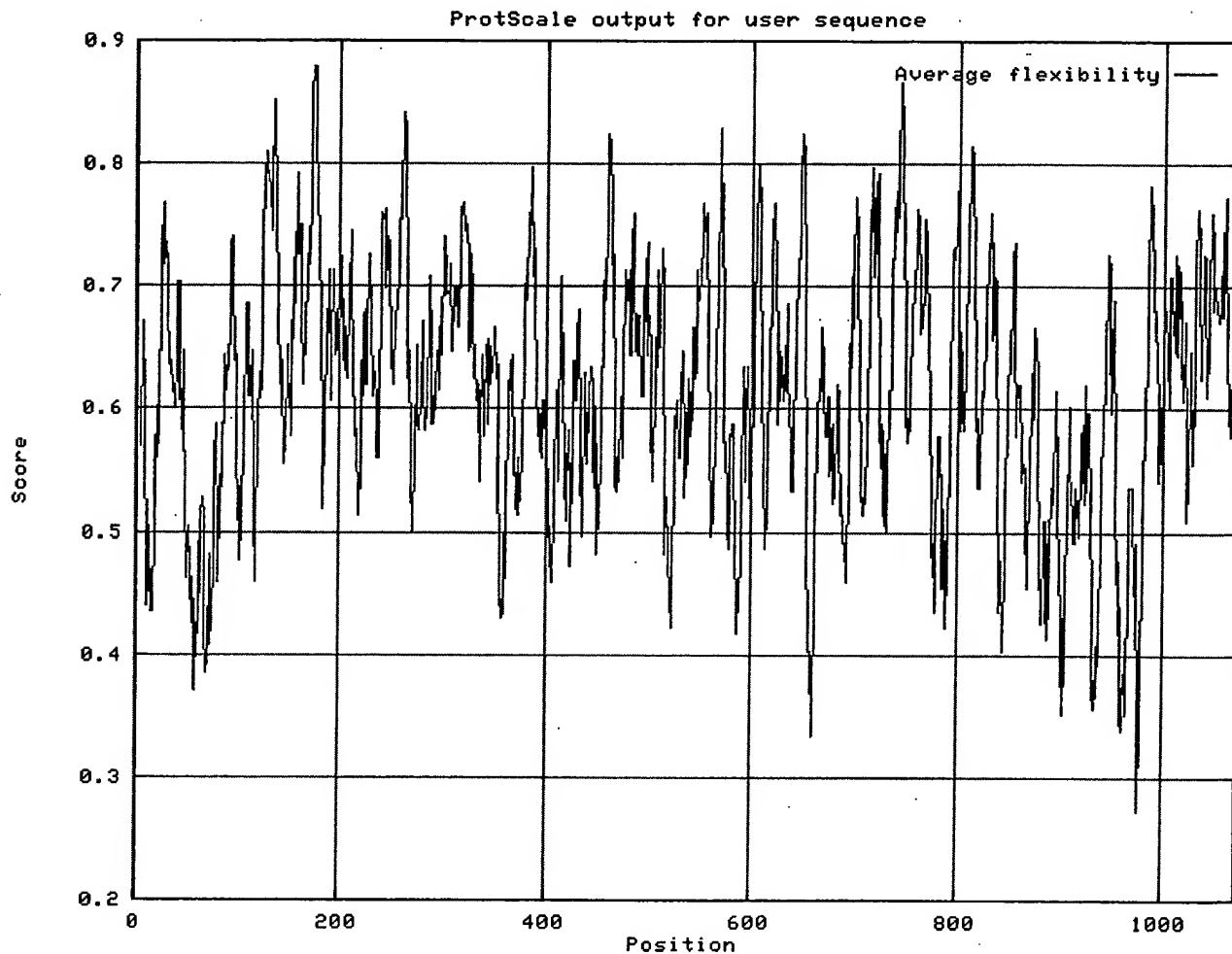
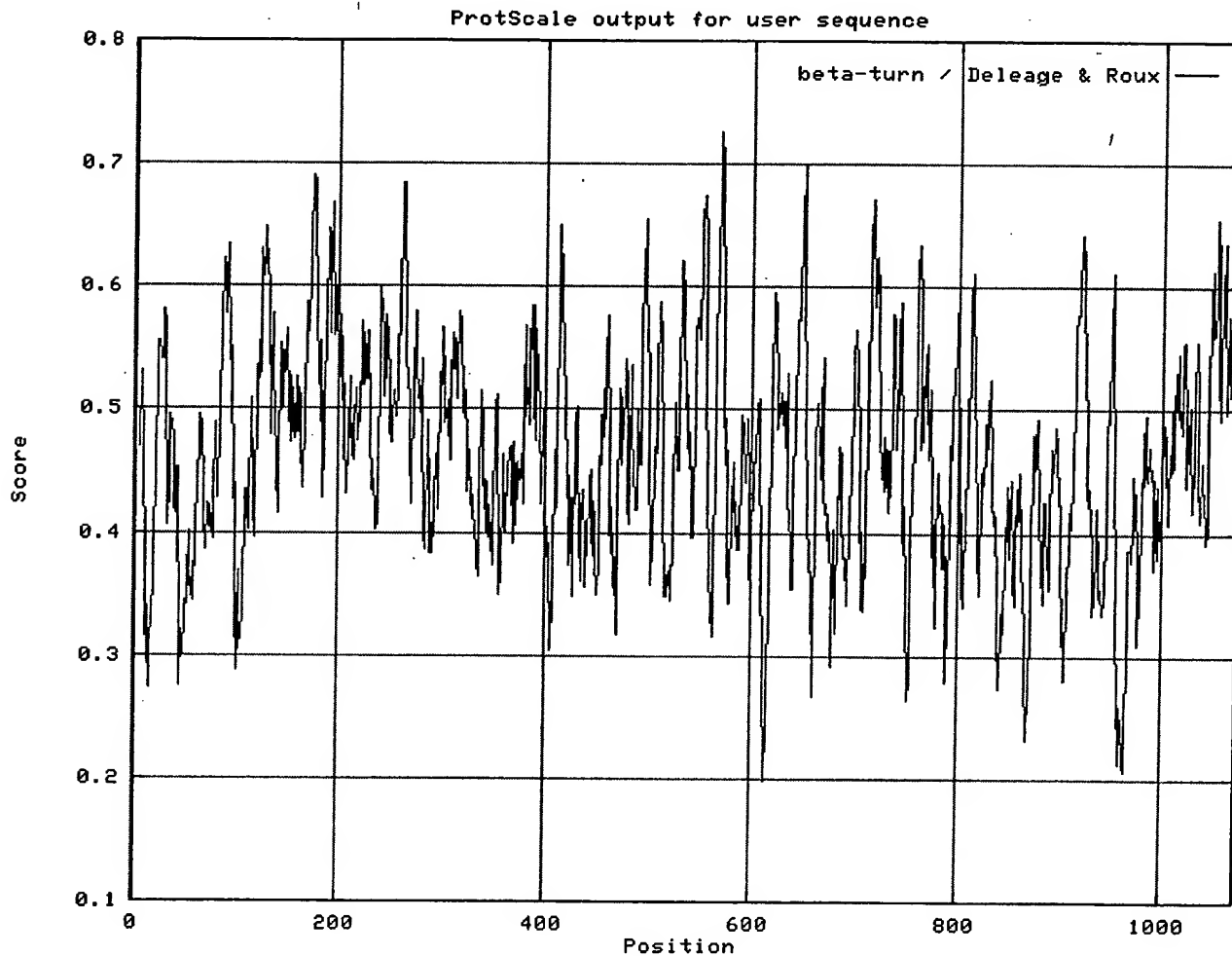
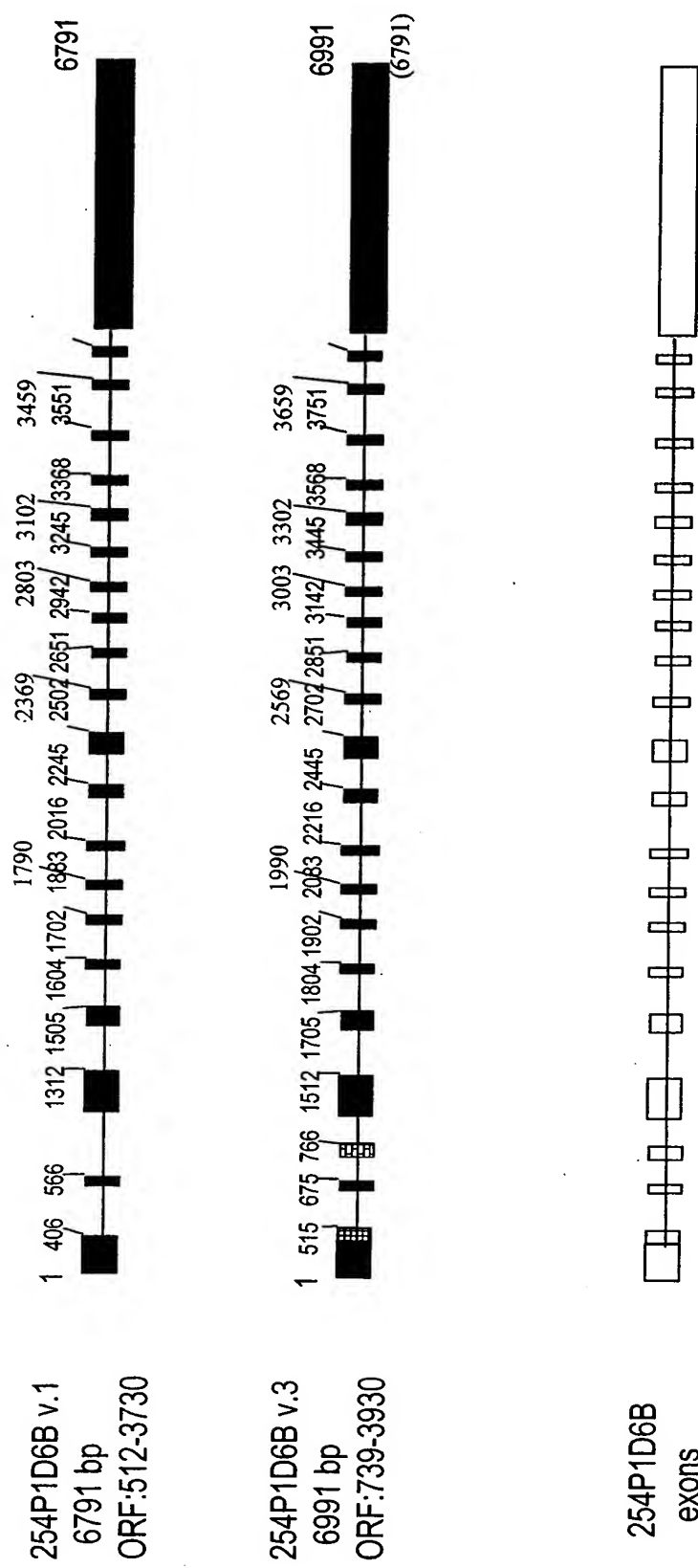


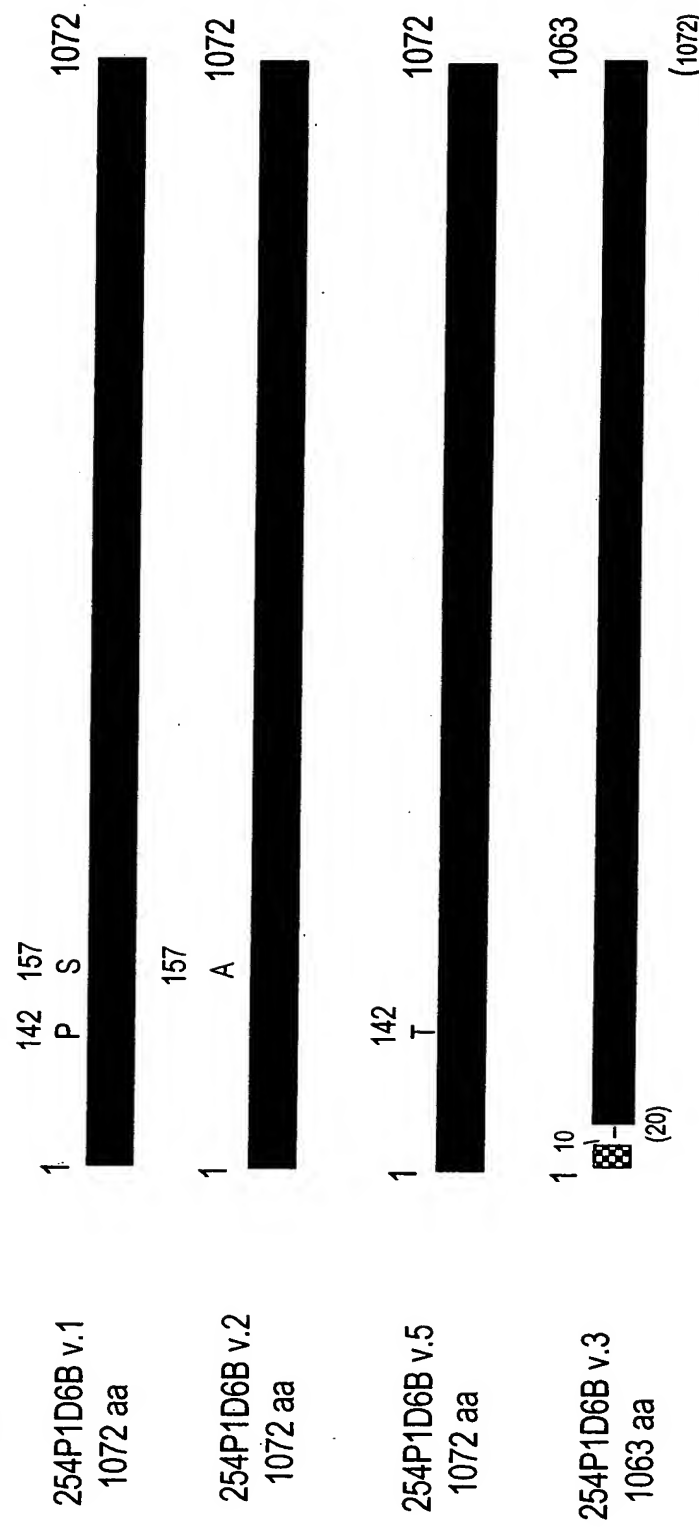
Figure 9: 254P1D6B variant 1  
Beta-turn Profile  
(Deleage, G., Roux B. 1987.  
Protein Engineering 1:289-294)



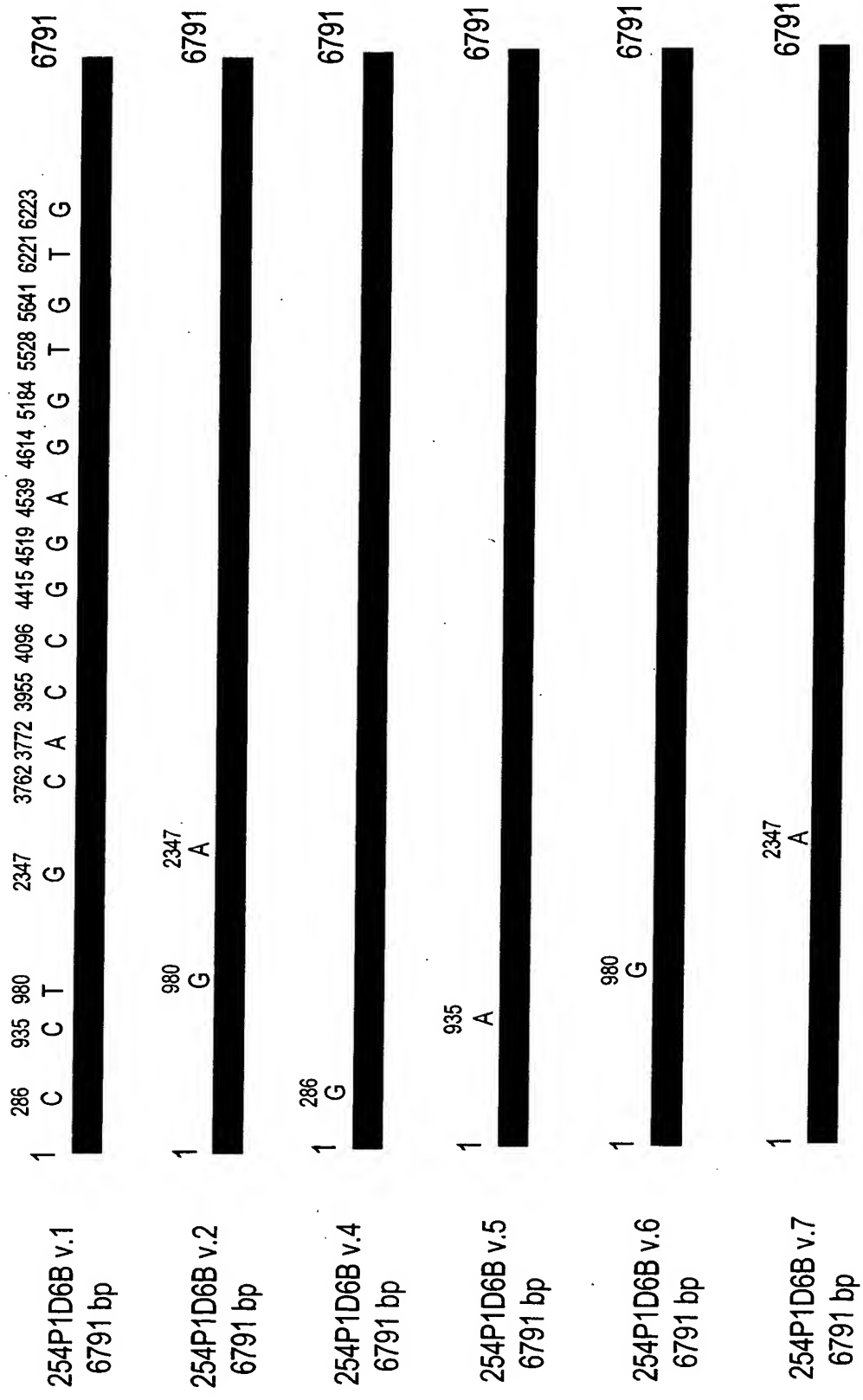
**Figure 10**



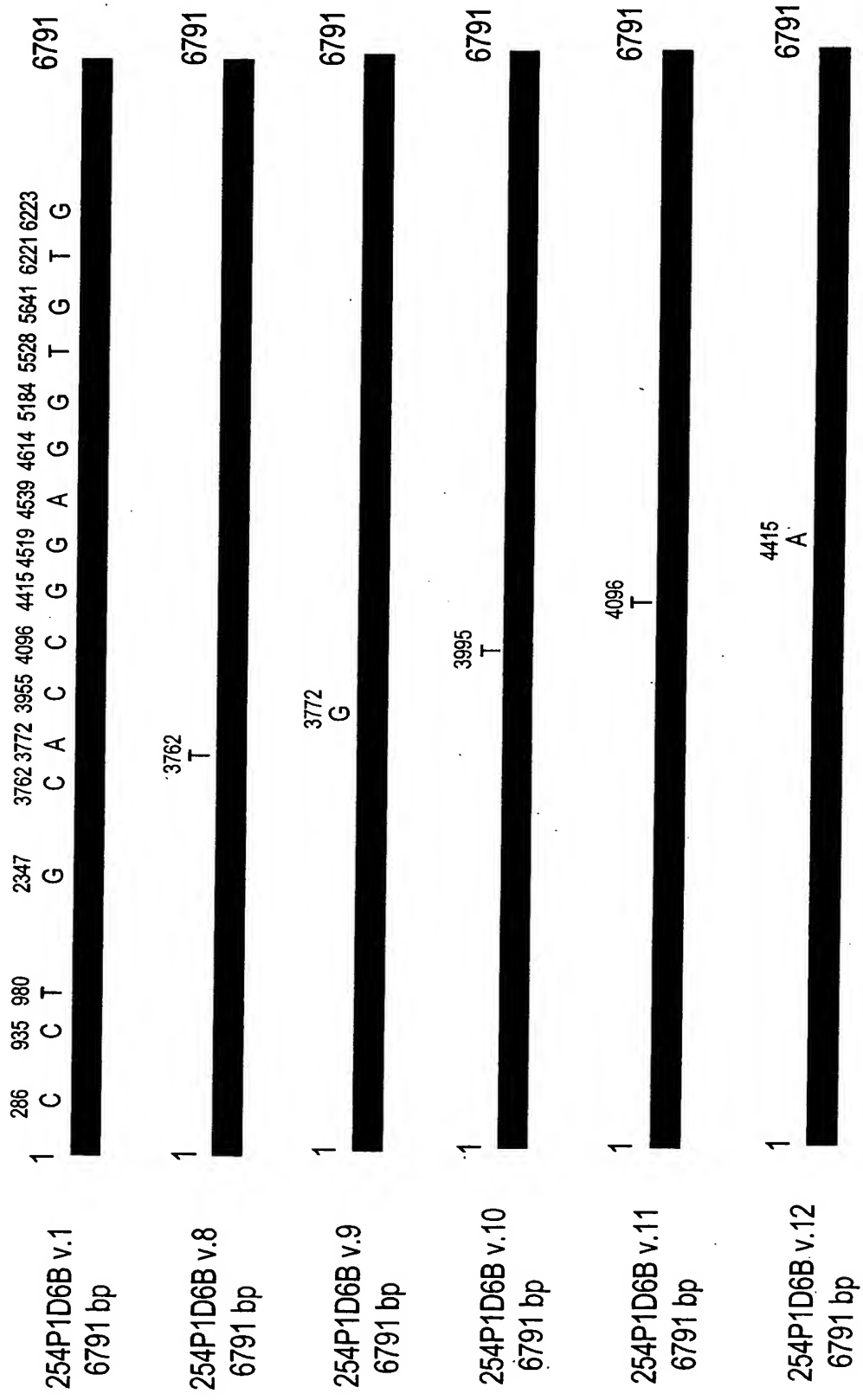
**Figure 11**

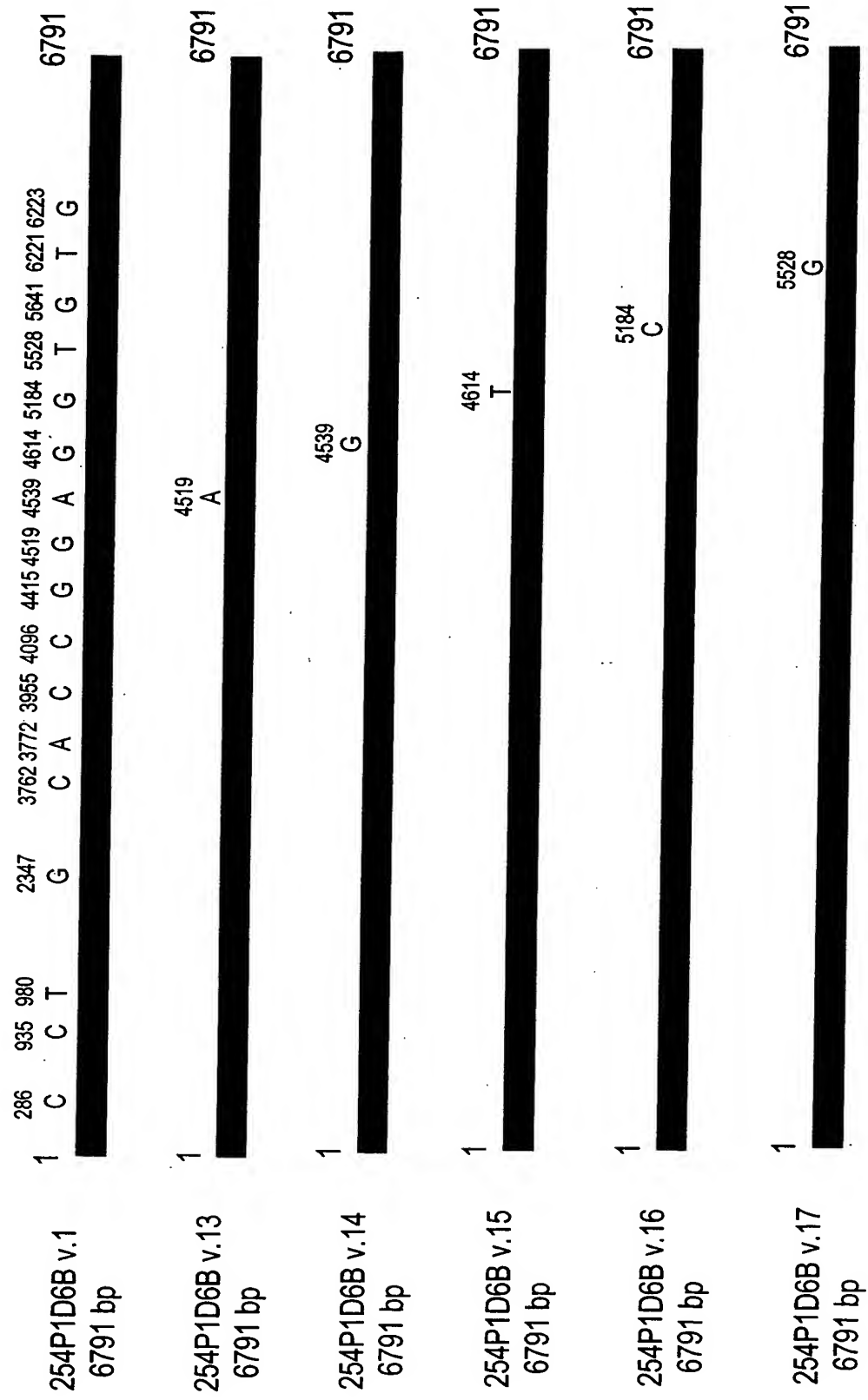


**Figure 12**



**Figure 12 (con'd)**



**Figure 12 (con'd)**

**Figure 12 (con'd)**

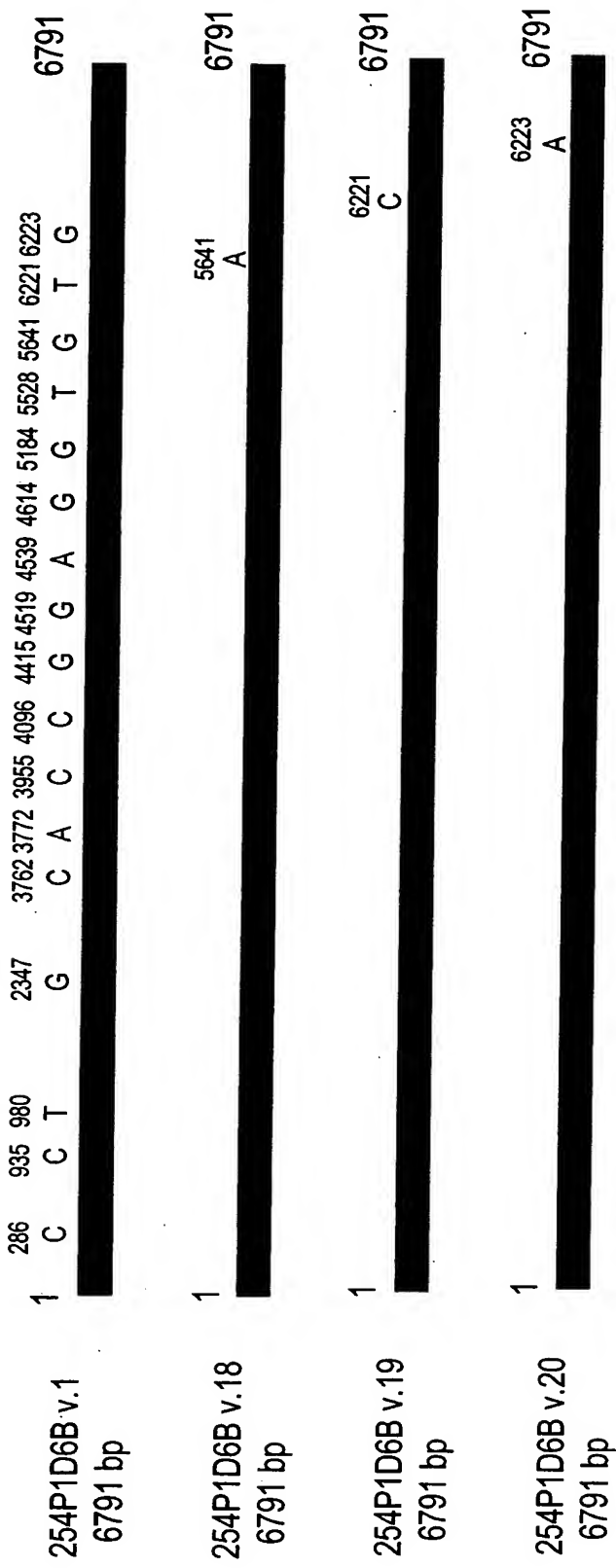
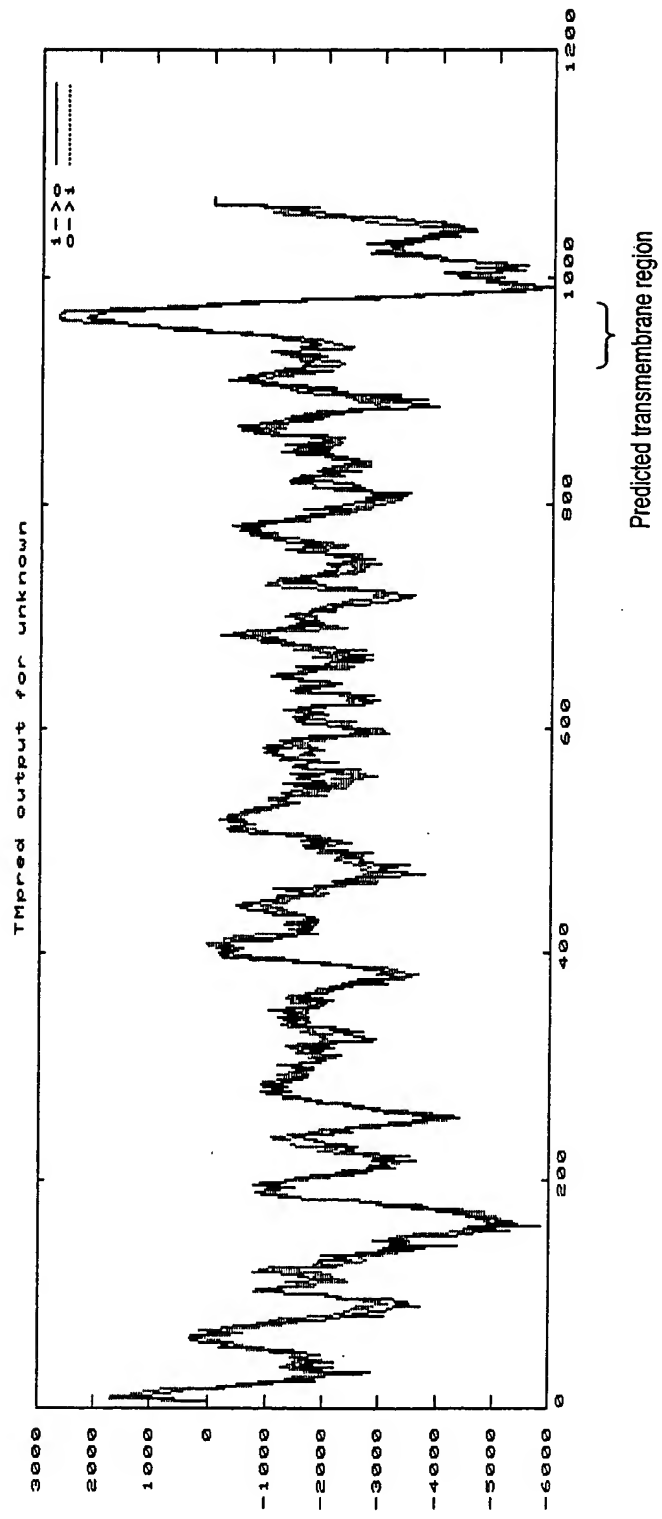


Figure 13: Secondary structure prediction of 254P1D6B variant 1

# Secondary structure prediction of 254P1D6B variant 1 (continued)

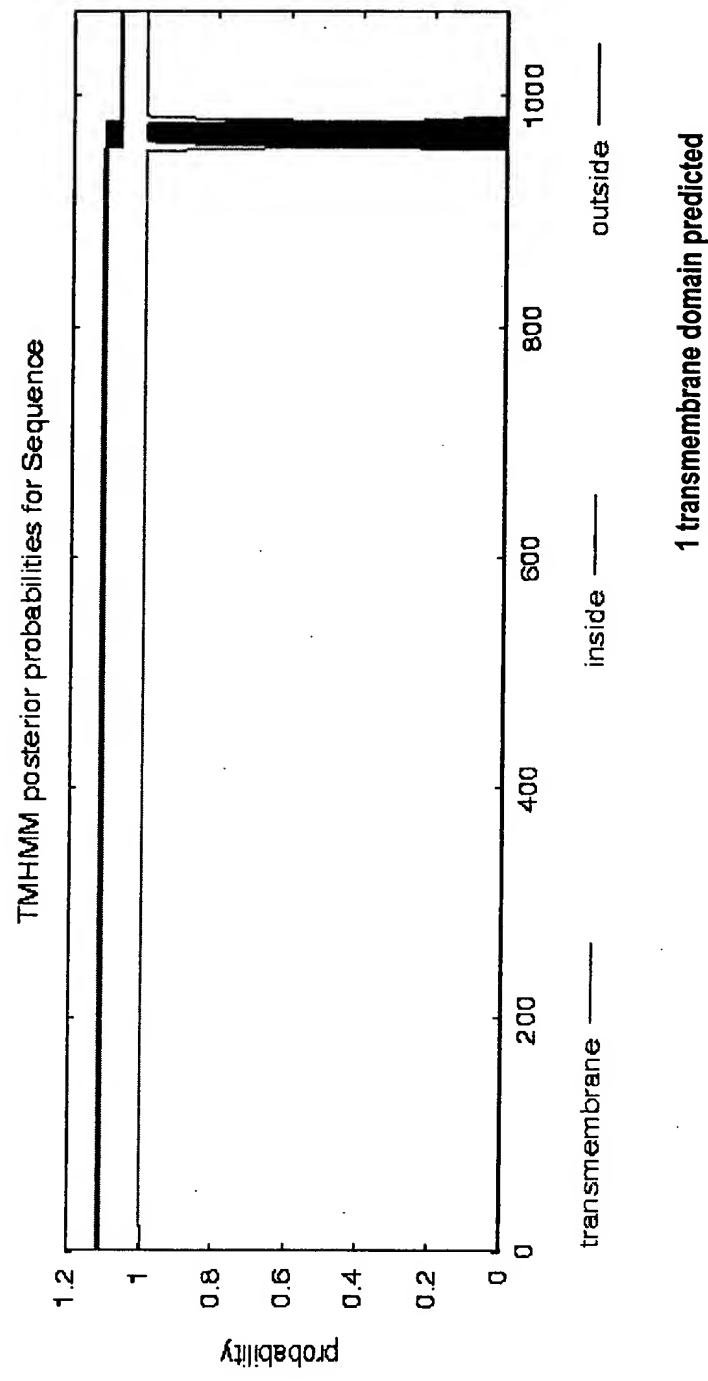
Alpha helix(h) : 18.19%  
 Extended strand (e) : 24.81%  
 Random coil(c) : 57.00%

**Figure: Transmembrane prediction for 254P1D6B variant 1  
13B**



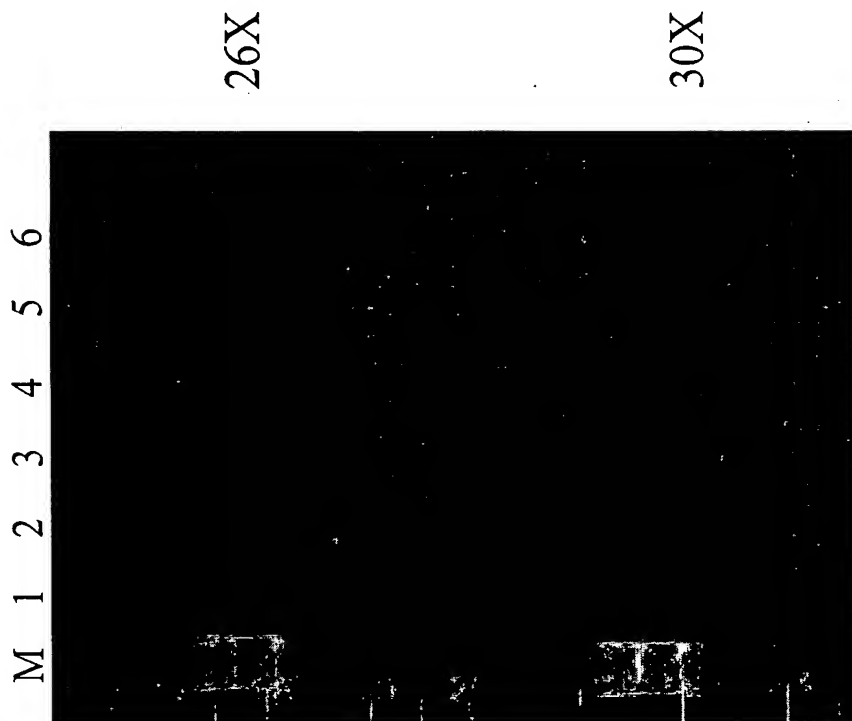
1 transmembrane domain predicted

Figure: Transmembrane prediction for 254P1D6B variant 1  
13C



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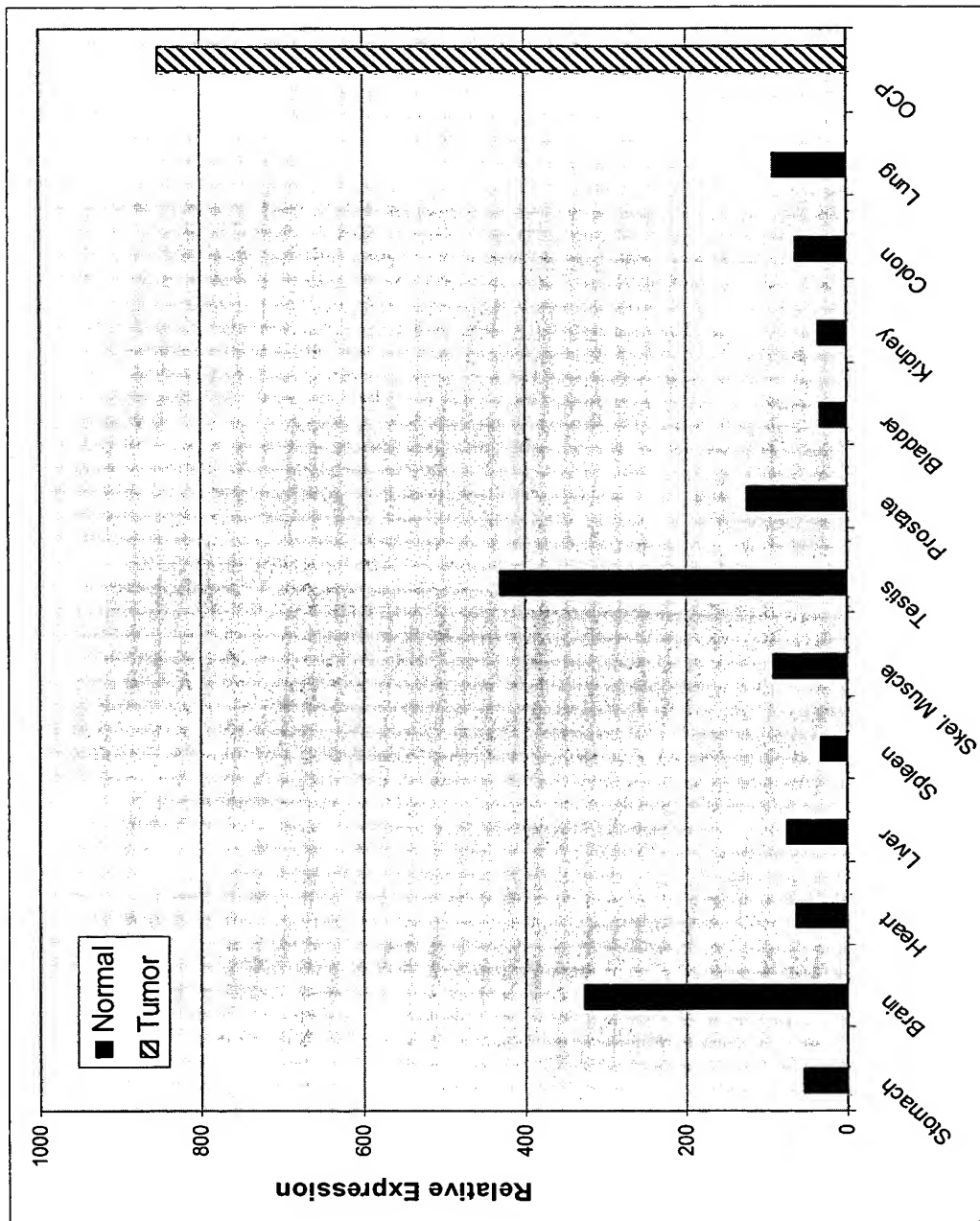
**Figure 14A 254P1D6B Expression by RT-PCR**

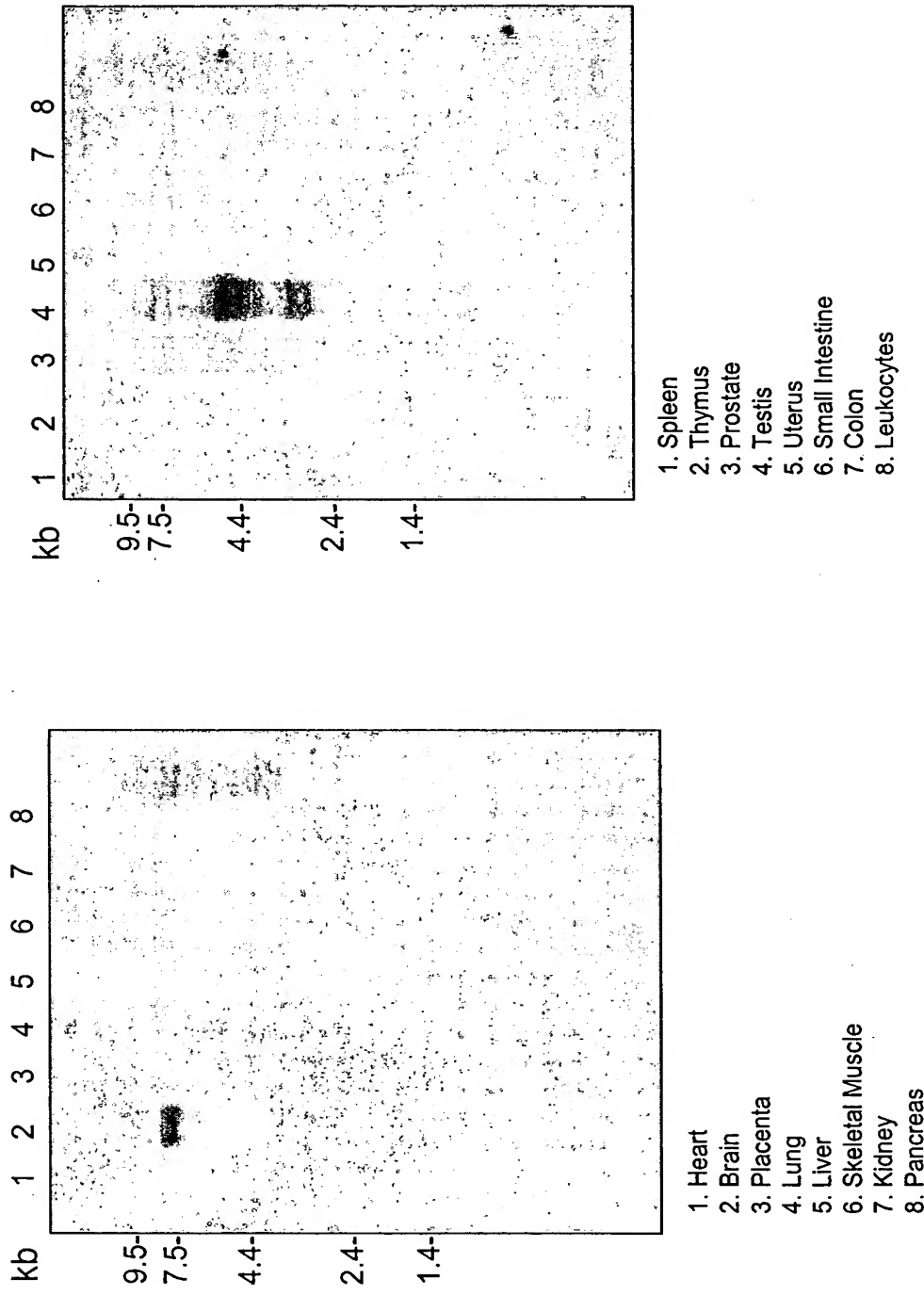


M = Marker

- 1) Vital Pool 1  
(Kidney, Liver, Lung)
- 2) Vital Pool 2  
(Colon, Pancreas, Stomach)
- 3) Normal Lung
- 4) Lung Cancer Pool
- 5) Ovary Cancer Pool
- 5) Pancreas cancer Pool

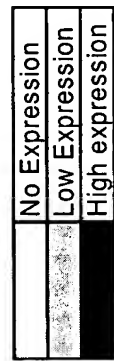
**Figure 14B Expression of 254P1D6B in  
Normal Human Tissues and Ovarian Cancer Patient Specimens**



**THIS PAGE BLANK****BEST AVAILABLE COPY****Figure 15 Expression of 254P1D6B in Normal Tissues**

**Figure 16 Expression of 254P1D6B in Lung Cancer Patient Specimens**

Panel#	Pathology	Grade	Expression
1	Normal		
2	A427 Cell line	3	
3	Adeno	I	
4	Adeno	IB	
5	Adeno	IIA	
6	Adeno	IIIA	
7	Adeno	IIIA	
8	Adeno	Mod Diff	
9	Adeno	Mod Diff	
10	Adeno		
11	Adeno		
12	Bronchioalv.	IA	
13	Large Cell	I	
14	Large Cell	IB	
15	Large Cell	IIA	
16	Large Cell	IV	
17	Papillary	I	
18	Papillary	IB	
19	Papillary	IV	
20	Small Cell		
21	Small Cell		
22	Small Cell		
23	Small Cell	IB	
24	Squamous	IB	
25	Squamous	IB	
26	Squamous	IB	
27	Squamous	IB	
28	Squamous	IB	
29	Squamous	IIA	
30	Squamous	IIA	
31	Squamous		
32	Squamous		
33	Squamous		



**Figure 17: Expression of 254P1D6b in 293T cells**

Figure  
17A  
293T-Vector  
293T-AGSx  
293T-254P1D6

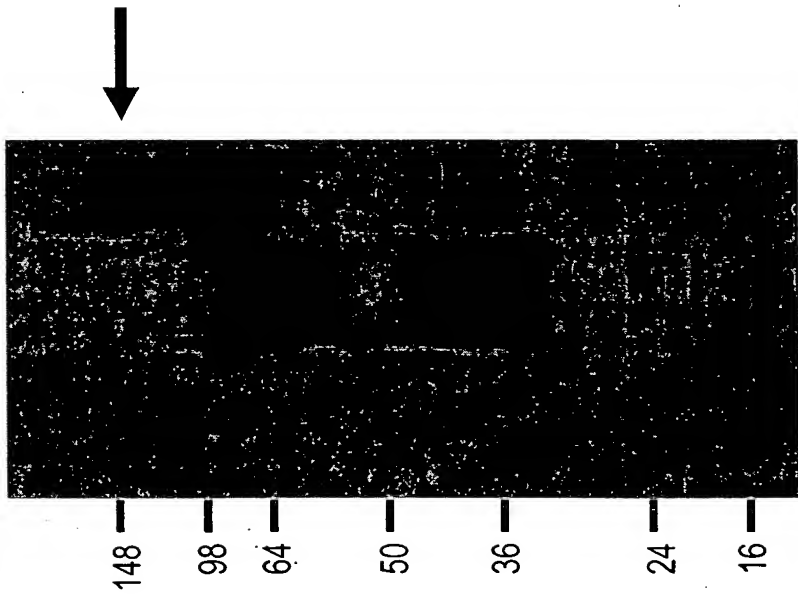
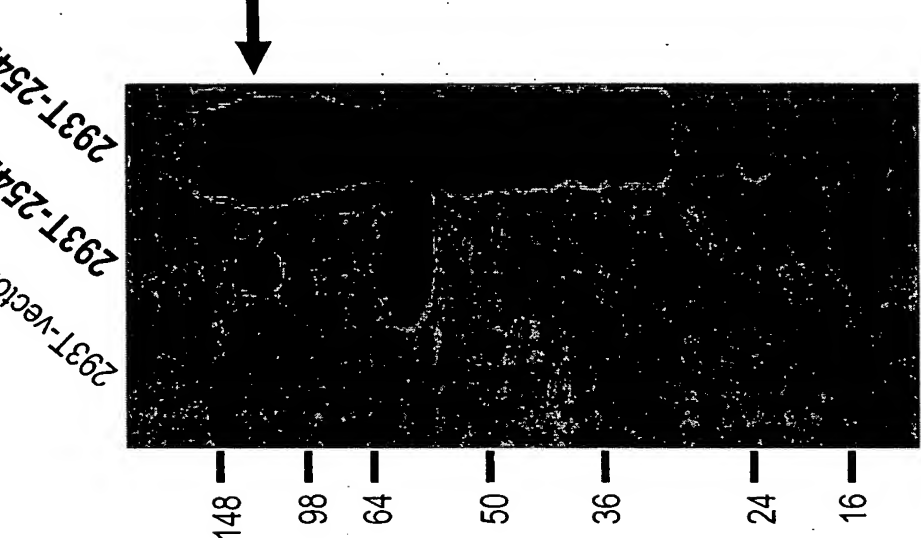


Figure  
17B  
293T-Vector  
293T-254P1D6  
293T-254P1D6  
media lysate



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